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(57) Abstract: The invention relates to a method of identifying plant-specific polypeptides and nucleic acids encoding them which are suitable as sites of action for finding herbicides, to the use of the polypeptides identified for identifying new, herbicidally active compounds, and methods of finding modulators of these polypeptides. Likewise, the invention relates to the use of the polypeptides in assay methods for identifying herbicidally active compounds.

-1-

# Polypeptides for identifying new herbicidally active compounds

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The invention relates to a method of identifying plant-specific polypeptides and nucleic acids encoding them which are suitable as sites of action for finding herbicides, to the use of the polypeptides which have been identified for identifying new herbicidally active compounds, and to methods of finding modulators of this polypeptide. Equally, the invention relates to the use of the plant polypeptides in assay methods for identifying herbicidally active compounds.

Herbicides have great importance in agriculture to avoid undesired plant growth by using herbicides. In modern agriculture, the use of herbicides constitutes an imperative factor for safeguarding yields and profits. This is where herbicides must meet increasingly high demands with regard to their efficacy, costs and above all their ecofriendliness. There is therefore a constant demand for new substances, known as lead structures, which can be developed into even more potent and even more ecofriendly new herbicides.

To date, only a few molecular sites of action, known as targets, play a key role for the action of herbicidal compounds. Three quarters of the entire herbicide market are dominated by just 5 targets, which are the sites of action of these herbicides: acetolactate synthase. elongases long-chain fatty for very acids. enolpyruvylshikimate-3-phosphate synthase, the photosystem  $\Pi$  and the auxin signal cascade. The remaining quarter of the market comprises just 6 further important targets: acetal-coenzyme A carboxylase, glutamine synthase, photosystem I, phytoene desaturase, protoporphyrinogen oxidase and tubulin. Herbicides for all of these targets have been known for over 20 years. During this period, herbicides with other, new targets have not gained market relevance. This situation leads to a thorough knowledge and exploitation of these targets in the search for new herbicidally active lead structures. At the same time, however, the use of new targets is extremely important for an innovation in the search for new lead structures for the development of novel and superior herbicides.

-2-.

To date it is generally customary to search for new lead structures in greenhouse tests. However, such tests require a good deal of labour and are expensive. The number of the substances, which can be tested in the greenhouse, is accordingly limited. However, even after suitable automation for increasing the throughput, greenhouse screening does not allow any findings as to whether substances may be directed against a new target. This must be determined in very complex subsequent experiments.

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An alternative to the search for lead structures which is nowadays generally customary is what is known as high-throughput screening or ultra-high-throughput screening (HTS or UHTS). This method, which was first established in pharmaceutical research, makes possible the automation of in-vitro assays for the search for lead structures for given targets. At the same time, it has been made possible to provide a high number of test substances by methods such as, for example, combinatorial chemistry. Thus, a multiplicity of methods has been developed as to how specific targets can be assayed by (U)HTS. The target-based search for lead structures for agricultural applications with the aid of (U)HTS does not differ from that for pharmaceutical applications and is therefore firmly established at present.

(U)HTS makes it possible to test the action of several hundreds of thousands of substances on a specific target within a few days. However, existing experience in industry shows that it is not possible to find a lead structure for each new target, at least not at present. It is therefore necessary to test a multiplicity of targets in order to identify suitable targets in addition to new herbicidal substances.

All of the five abovementioned herbicide targets which dominate the market, and most of the remaining targets, are only found in plants but not in animals. This is no coincidence but is due to the advantageous properties of such active compounds. Thus, there is only little danger of a toxic effect on humans and the environment in

plant-specific targets. This can be proved by comparing the two targets acetolactate synthase and protoporphyinogen oxidase. At the beginning of the 80s, highly effective and innovative compounds were discovered for both targets, initially without knowing the target. A series of herbicides were quick to reach the market in the case of the plant-specific target acetolactate synthase, so that acetolactate synthase is currently ranked third among the herbicide targets. Even though a very large variety of herbicides which act on protoporphyrinogen oxidase, which is also found in animals, is now known, the unfavourable toxicology of these products has as yet not led to an important commercial product.

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Toxicological studies are complicated and expensive. As a rule, these studies are only performed when a certain basic development of new lead structures has already taken place. Even so, the research expenses up to this point are quite considerable. It is therefore advantageous to minimize the toxic effect of new herbicides, which is due to the target, right at the beginning. This can be achieved by simply using those targets for the search for lead structures which are found only in plants, but not in animals.

Especially advantageous targets for new herbicides are searched for in essential biosynthetic pathways. Thus, for example, the biosynthesis of isoprenoids, building blocks of carotinoids and of plastoquinone and chlorophyll, are imperative for the growth of plants. The inhibition of a step in this plant-specific biosynthetic pathway, also known as the 1-desoxyxylulose-5-phosphate pathway, leads to the death of a plant (DE 199 35 967). The knowledge of the plant specificity of specific metabolic pathways is currently fundamental knowledge in plant biochemistry (see, for example, B. B. Buchanan, W. Gruissem and R. L. Jones (Editors); "Biochemistry & Molecular Biology of Plants", American Society of Plant Physiologists, Rockville, MD, USA; 2000), even when it remains partially unclear which role certain proteins take on in the plant, and whether corresponding proteins or those with an equivalent task are also found in, for example, mammals.

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Each new candidate herbicide must meet a number of criteria before it can be approved, and the choice of a suitable target is the first step in this search.

It is helpful to consider the existing genome information which is now available to the public, and to take note of some key criteria of herbicidal active compounds:

- 1. An active compound should be sufficiently selective and produce a herbicide which should be specific, or at least very selective, for plants (with regard to humans or animals).
- 2. An active compound should attack proteins or else genes which are imperative for the growth or the viability of the undesired plants, and
- 3. something should be known about the function of the target protein or target gene so that an assay and high-throughput screens can be established.

It is furthermore important for choosing suitable targets that the probability of identifying a new lead structure is considerably higher when the target has a natural binding property for ligands of low molecular weight. This is in contrast to, for example, individual protein components of large complexes with many subunits. The interference of protein-protein interactions by small ligands is less possible and requires, in principle, larger active compounds whose production costs are then frequently higher, so that a meaningful use of these active compounds as herbicides is made substantially more difficult. Targets with small natural ligands are, for example, enzymes, receptors and channels. Moreover, enzymes, receptors and channels can frequently be assayed more easily in assay methods (HTS or UHTS) than other proteins.

A possibility of recognizing plant-specific new targets is to test the enzymes or receptors and channels involved in plant-specific metabolic pathways or signal chains one after the other, using present-day biochemical knowledge (B. B. Buchanan, W.

Gruissem and R. L. Jones (Editors); "Biochemistry & Molecular Biology of Plants", American Society of Plant Physiologists, Rockville, MD, USA; 2000). However, this route carries the risk of overlooking important properties of the proteins.

While new routes for, for example, based on sequence information in the field of antibiotic research have already been described (see, for example, Molly B. Schmid, Novel approaches to the discovery of antimicrobial agents, Curr. Opin. Chemical Biol., 2, 529-534, 1998.), a method of identifying suitable targets for the search for herbicides on the basis of existing data from sequencing work is as yet not available.

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It was therefore the object of the present invention to describe a method which is suitable for identifying, in an efficient and reliable fashion, those nucleic acids or polypeptides encoded by them from among sequence information available in public databases, which can be used for the search for new herbicidal active compounds as plant-specific sites of action which can be obtained by a screening method. The object of the present invention was also to identify and to describe suitable target proteins by means of the method described and to make these available for use in screening methods for the search for new active compounds.

The complete knowledge of the genome of *Arabidopsis*, of humans and of many other organisms now allows to filter out, by means of computer-aided comparison of the proteins encoded in the genome, those proteins which occur in one organism but not another. Thus, it is also possible to recognize plant-specific proteins whose

function was hitherto unelucidated.

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In the present context, the term "plant-specific" is understood as meaning that no similarity with proteins from animals, in particular higher animals (Metazoa; in particular Chordata) is found.

A series of these plant-specific proteins, however, are also found in micro-organisms (for example bacteria, fungi).

- 6 -

In the present invention there is now described a possibility of identifying, from publicly available information and with the aid of computer-aided methods, those proteins and the nucleic acids encoding them which are suitable for use in methods for identifying new herbicidally active compounds.

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The comparison of the proteins encoded in various genomes is possible by means of a systematic alignment comparison (for example BLAST (Altschul et al., 1990), FastA (Lipman and Pearson, 1985, Pearson 1991), Search (Smith and Waterman, 1981) Hmmer (Durbin et al., 1998)) between all proteins of one organism and those of the other organism. Preferably, one organism is selected, and the presence of the homologous sequence in other organisms is then studied.

In the present invention, all of the proteins encoded in the genome of *Arabidopsis* thaliana (hereinbelow abbreviated to "Arabidopsis") are compared with all of the other sequences which are accessible in public databases. The following databases were used as source for the Arabidopsis polypeptides in the present invention:

- a) TAIR (Huala et al., 2001), which is a searchable relational database comprising information related to *Arabidopsis thaliana*, and
- b) GenBank (Benson et al., 2000), which is the NIH genetic sequence database, an annotated collection of all publicly available DNA sequences, including protein translations.
- Databases which can be used for the comparison are, for example, the following:
  - a) SwissProt, which is a curated protein sequence database and provides a high level of annotations (e.g. function, domains structure, variants etc.)
- b) TrEMBL and TrEMBL-New (non-redundant protein databases), which are computer-annotated supplements of Swiss Prot and contain all the translations of EMBL nucleotide sequence entries not yet integrated in SwissProt and

-7-

whereby TrEMBL-New is a weekly update to TrEMBL which contains the protein-coding sequences from EMBLNEW

PCT/EP01/09892

(see Bairoch and Apweiler, 2000).

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WO 02/10210

All of the protein-encoding genes, and/or the polypeptides encoded by them, of the databases are compared with each other (pair-wise comparison; each polypeptide with each polypeptide) in order to find homologous similarities. The rigorous Smith-Waterman algorithm is used for this purpose.

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To assess whether a given alignment constitutes evidence for homology, it helps to know how strong an alignment can be expected from chance alone. A local alignment without gaps consists simply of a pair of equal length segments, one from each of the two sequences being compared. A modification of the Smith-Waterman or Sellers algorithms will find all segment pairs whose "scores" can not be improved by extension or trimming. These are called high-scoring segment pairs (HSPs). To analyze how high a score is likely to arise by chance, a model of random sequences is needed. For proteins, the simplest model chooses the amino acid residues in a sequence independently, with specific background probabilities for the various residues. In the limit of sufficiently large sequence lengths m and n, the statistics of HSP scores are characterized by two parameters, K and lambda. Most simply, the expected number of HSPs with score at least S is given by the formula

## $E = Kmne^{-\lambda S}$

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which is the so called E-value for the score S. The parameters K and lambda can be thought of simply as natural scales for the search space size and the scoring system respectively.

The measure for the similarity which is obtained is therefore an E-value (expect-value). As shown above, the E-value indicates the probability of which the existing

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agreement between two proteins or else genes or nucleic acids is due to pure random chance. The smaller the E-value, the more significant a hit in the search. If, for example, the E-values are in the range of 1e-70, this means that owing to the size of the database, only  $10^{-70}$  hits would have been expected with the search sequence. This also means that the results are highly significant. In the case of two identical sequences, the E-value thus progresses towards zero. In the case of two entirely unrelated sequences, the E-value converges to values greater than one.

In the present method according to the invention, the criterion chosen for plant specificity and thus the suitability of the polypeptide according to the present invention, the E-value was chosen such that the exponent of the E-value of a paralogous or orthologous plant amino acid sequence must exceed that of a corresponding paralogous or orthologous animal or human sequence, in as far as such an animal or human sequence exists, at least by a factor of 3. The E-value of 10<sup>-30</sup> is particularly suitable as limit for defining plant specificity. If the abovementioned factor decreases, it can be assumed with high probability that the homology between the plant sequence and the animal or human sequence is too high to classify a plant polypeptide as plant-specific and suitable for the use according to the invention in methods of finding herbicides.

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The term "identity" as used in the present context refers to the number of sequence positions which are identical in an alignment. In most cases, it is indicated as a percentage of the alignment length.

The term "similarity" as used in the present context, in contrast, requires the definition of a similarity matrix, that is to say a measure for the degree of similarity one wishes to assume between, for example, a valine and a threonine or a leucine.

The term "homology" as used in the present context, in turn, refers to evolutionary relationship. Two homologous proteins have developed from a joint precursor sequence. The term does not necessarily imply identity or similarity, apart from the

-9-

fact that homologous sequences are usually more similar (or have more identical positions in an alignment) than non-homologous sequences.

The term "orthologues" or "orthologous" as used in the present contexts refers to a functional counterpart, for example a protein in another organism, both having developed from a shared precursor. Normally, orthologues retain a shared function. In contrast, "paralogues" are genes or proteins resulting therefrom which have originated by duplication within a genome and which have assumed different functions during evolution which may still have similarity with each other.

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Proteins are termed orthologous when

1. they have the highest level of pair-wise similarity (compared with the identities of the two proteins with all the other proteins in other genomes) and

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2. the similarity is significant (E<0.01).

The proteins encoded in the *Arabidopsis* genome and the results of the comparison with all the other public sequences were stored in a relational database (Oracle) in the present invention.

Such a relational database model was presented in 1970 by Codd et al. All of the data to be processed are shown in Tables (relatins) with a fixed number of columns and any desired number of lines (tupels). Data redundancies are avoided by distributing the information to individual tables. To date, this model remains the basis of most of the commercial database systems.

In general, the assigning of a description which is firstly correct and can secondly be searched for readily, what is known as an annotation, to each sequence constitutes a major problem in practice. An "annotation" of a sequence is the assigning of biologically relevant properties to this sequence of parts thereof.

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By comparison of, possibly competing, alternative annotations in public databases and by individual corrections, a standardized annotation for each database entry has now been generated in the present invention. For example, the annotation takes such a form that the description of enzymes, receptors and channels (transporters) starts with the respective functional name, that is, for example, with "acetolactate synthase".

An annotation was assigned to the sequence in a multi-step process: first, the information content of words or terms within a sequence description were analysed and these words/terms were correspond categorized. Thus, the description "acetolactate synthase" leads to more information on a sequence than the descriptions "Unknown Protein" or "Hypothetical Protein" or "exon predicted by xgrail, quality marginal\_shadowexon". This procedure first gives two categories of words/terms and, based on these categories, eventually two categories or sequence descriptions: those with a low information content and those with a high information content.

Only the sequence descriptions with a high information content are used for assigning an annotation to a sequence. These annotations obtained in this way are subsequently aligned in a suitable fashion with the annotations obtainable from TAIR. In the present invention, the TAIR annotation for a given sequence was adopted if such an annotation did exist.

This process was automated by developing suitable programs.

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In a final step, the present annotations were rechecked and, if appropriate, corrected, to arrive at the final standardized annotation.

The database established within the present invention contains sequences from Arabidopsis and the relevant descriptions (annotations) and E values in question and thus makes possible an efficient and meaningful analysis of the sequence data, which

- 11 -

results in the reliable identification of suitable plant-specific targets for the purposes of the present invention.

All the enzymes, receptors and channels or transporters with the above-described plant-specific E-values were then filtered out from the annotations of the database according to the invention with the aid of a suitable algorithm with suitable search terms. The polypeptides found by this method are shown in Table 1. In addition to the annotation of the polypeptide whose sequence is available by means of the reference to the sequence listing in the present application, Table 1 also shows which particular class of polypeptides it belongs to. Enzymes were arranged for example by classes such as "dehydrogenase" or "oxygenase". Receptors were searched for with the search term "receptor", but not "receptor kinase". Channels were searched for with the search term "channel" or "transporter". The table also contains what is known as the accession number of the sequence, in as far as it is known. The accession number provides information on the database or the number in which, or under which, the polypeptide sequence in question can be found. Furthermore, the table contains references to known homologous sequences from other organisms and a reference to the SEQ ID NO. under which the sequence in question is filed in the sequence listing.

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## Table 1:

ENTRY	SEQ	DESCRIPTION	CLASS-
ID NO	ID NO		
5	1	INORGANIC PYROPHOSPHATASE, PUTATIVE SIMILAR	Phosphatase
		TO SOLUBLE INORGANIC PYROPHOSPHATASE GB:	
		AAD46520 GI:5669924 FROM [POPULUS TREMULA X	
		POPULUS TREMULOIDES]	
12	2	FATTY ACID ELONGASE 3-KETOACYL-COA SYNTHASE	Synthase
		1 IDENTICAL TO GB:AAC99312 GI:4091810 FROM	
		[ARABIDOPSIS THALIANA]	
33	3	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL, PUTATIVE SIMILAR TO CYCLIC	
]		NUCLEOTIDE AND CALMODULIN-REGULATED ION	
		CHANNEL GB:CAB40128 GI:4581201 FROM	
ļ		[ARABIDOPSIS THALIANA]	
38	4	FLAVONOL 3-O-GLUCOSYLTRANSFERASE, PUTATIVE	Transferases
		SIMILAR TO FLAVONOL 3-O-GLUCOSYLTRANSFERASE	
		GB:Q40287 FROM [MANIHOT ESCULENTA]	
41	5	FLAVONOL 3-O-GLUCOSYLTRANSFERASE, PUTATIVE	Transferases
		SIMILAR TO FLAVONOL 3-O-GLUCOSYLTRANSFERASE	
		GB:Q40287 FROM [MANIHOT ESCULENTA]	
46	6		Kinase
		KINASE(ATPIP5K1) GI:3702691 FROM [ARABIDOPSIS	
		THALIANA][HYPOTHETICAL PROTEIN CONTAINS	
		SIMILARITY TO]	
50	7	DEHYDROGENASE GI:1922246 FROM [ARABIDOPSIS	Dehydrogenases
		THALIANA] UNKNOWN PROTEIN SIMILAR TO	
		PUTATIVE	<u> </u>
53	8	SERINE/THREONINE KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		SERINE/THREONINE KINASE GI:7248457 FROM	ļ
		[LOPHOPYRUM ELONGATUM]	Deduction
57	9	FERRIC REDUCTASE LIKE TRANSMEMBRANE	Keduciase
	10	COMPONENT	D. J. J.
58	10	FERRIC REDUCTASE LIKE TRANSMEMBRANE	Keductase
		COMPONENT	

72	11	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN Kinase, Protein
, , , ,		KINASE GB:BAA24694 GI:2852447 FROM [ARABIDOPSIS]
		THALIANA]
77	12	K EFFLUX ANTIPORTER KEAI IDENTICAL TO Transporter
		GB:AAD01191 GI:4101473 FROM [ ARABIDOPSIS
		THALIANA]
83	13	DIMETHYLADENOSINE TRANSFERASE, PUTATIVE, 5'Transferases
		PARTIAL SIMILAR TO DIMETHYLADENOSINE
		TRANSFERASE GB:AAC09322 GI:3005590 FROM
		[ARABIDOPSIS THALIANA]
85	14	CUCUMISIN-LIKE SERINE PROTEASE GB:AAC18851 Protease
		GI:317687 FROM [ARABIDOPSIS
		THALIANA]4[HYPOTHETICAL PROTEIN CONTAINS
ļ		SIMILARITY TO]
87	15	RIBULOSE-1,5 BISPHOSPHATE CARBOXYLASE Carboxylase
		GB:L34291 GI:508550 FROM [PISUM
!		SATIVUM][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
93	16	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45849 Oxidase
1		GI:5262223 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]
95	17	NUCLEOTIDE SUGAR EPIMERASE, PUTATIVE SIMILAR Epimerase
		TO NUCLEOTIDE SUGAR EPIMERASE GB:AAC18831
		GI:3093975 FROM [VIBRIO VULNIFICUS]
100	18	CHALCONE SYNTHASE HOMOLOG, GP U90341 2507617 Synthase
100	10	AND ANTHER SPECIFIC PROTEIN, GP Y14507 2326772
118	10	SER-THR PROTEIN KINASE-LIKE PROTEIN GI:9294588 Kinase, Protein
] 118	19	
}		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL
		PROTEIN SIMILAR TO]
127	20	(1-4)-BETA-MANNAN ENDOHYDROLASE PRECURSOR, Hydrolase
		PUTATIVE SIMILAR TO (1-4)-BETA-MANNAN
		ENDOHYDROLASE PRECURSOR GI:9836826 FROM
		[LYCOPERSICON ESCULENTUM]
132	21	CHITINASE, PUTATIVE SIMILAR TO CHITINASE Chitinase
		GI:1237025 FROM [ARACHIS HYPOGAEA]
136	22	DIOXYGENASE, PUTATIVE SIMILAR TO DIOXYGENASE Oxygenases
		GI:1666096 FROM [MARAH MACROCARPUS]

142	23	POLYGALACTURONASE PG1, PUTATIVE SIMILAR TO Glycosylase	
		POLYGALACTURONASE PG1 GB:AAD46483 GI:5669846	
		FROM [GLYCINE MAX]	
154	24	ALDEHYDE OXIDASE, PUTATIVE SIMILAR TO	
		ALDEHYDE OXIDASE GB:BAA28630 GI:3172044 FROM	
		[ARABIDOPSIS THALIANA]	
157	25	NUCLEOSIDE TRANSPORTER GB:AAF26446 GI:6715514 Transporter	
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO][PUTATIVE]	
158	26	BETA-XYLOSIDASE, PUTATIVE SIMILAR TO BETA-Glycosylase	
		XYLOSIDASE GB:Z84377 GI:2102655 FROM	
		[ASPERGILLUS NIGER]	
166	27	GLYCOSYL TRANSFERASE GB:CAB80706 GI:7268597 Transferases	
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO][PUTATIVE]	
167	28	CELLULOSE SYNTHASE CATALYTIC SUBUNIT Synthase	
		GB:AAC39336 GI:2827143 FROM [ARABIDOPSIS]	
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]	
172	29	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase	
		POLYGALACTURONASE GI:288611 FROM [ZEA MAYS]	
174	30	PECTINESTERASE, PUTATIVE SIMILAR TO PECTIN Esterase	į
		ESTERASE GI:1213628 FROM [PRUNUS PERSICA]	
185	31	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Transferases	
		TO GLUTATHIONE S-TRANSFERASE GI:860955 FROM	
196		[HYOSCYAMUS MUTICUS]	
186	32	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Transferases TO GLUTATHIONE S-TRANSFERASE GI:860955 FROM	
		[HYOSCYAMUS MUTICUS]	
187	33	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Transferases	
10/	55	TO GLUTATHIONE S-TRANSFERASE GI:860955 FROM	
		[HYOSCYAMUS MUTICUS]	
188	34	GLUTATHIONE-S-TRANSFERASE, PUTATIVE SIMILAR Transferases	
		TO GLUTATHIONE-S-TRANSFERASE GI:169887 FROM	
		[SILENE VULGARIS]	
L			

197	35	LIPOAMIDE DEHYDROGENASE COMPONENT OF THE	Dehydrogenases
		PYRUVATE DEHYDROGENASE COMPLEX E3,	
		CONTAINS PF 00010 HELIX-LOOP-HELIX DNA-BINDING	
		DOMAIN. ESTS GB T45640 AND GB T22783 COME FROM	
		THIS GENE[PUTATIVE]	
209	36		Kinase
209	30	GENEFINDER	remaso
224	37	ISOAMYLASE SIMILAR TO GI 1652733 GLYCOGEN	Glycocylaca
224	)	OPERON PROTEIN GLGX FROM SYNECHOCYSTIS SP.	Gly cosy lase
		GENOME GB D90908. ESTS GB H36690, GB AA712462,	
		GB AA651230 AND GB N95932 COME FROM THIS	
		GENE[PUTATIVE]	
232	38	l	Transferases
	30	HYDROXYCINNAMOYL/BENZOYLTRANSFERASE[HYP	Transfords
<u> </u>		OTHETICAL PROTEIN SIMILAR TO	
233	30	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
255	37	SIMILAR TO ESTS EMB Z34690, GB T04168, GB H37738,	Oxiduse
		GB T76913, GB T43801, AMD GB T21964[PUTATIVE]	
234	40	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
25.		SIMILAR TO ESTS EMB Z34690, GB T04168, GB H37738,	
		GB T76913, GB T43801, AMD GB T21964[PUTATIVE]	
243	41	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE	Transferases
		GB:BAA74428[HYPOTHETICAL PROTEIN SIMILAR TO]	
250	42	PROTEIN PHOSPHATASE SIMILAR TO	Phosphatase
		GB:AAB97706[PUTATIVE]	-
254	43	PROTOCHLOROPHYLLIDE REDUCTASE SIMILAR TO	Reductase
		PROTOCHLOROPHYLLIDE REDUCTASE PRECUSOR;	į
		SIMILAR TO ESTS GB R30630, GB T46162, EMB Z26728,	İ
		GB AA042736, AND GB AA042730[PUTATIVE]	· [
284	44	ANTHOCYANIN 5-AROMATIC	Transferases
		ACYLTRANSFERASES[HYPOTHETICAL PROTEIN	·
		NEARLY IDENTICAL TO], ARABIDOPIS PROTEIN	
		F21B7.22, SIMILAR TO	ļ
294	45	ACID PHOSPHATASE; LOCATION OF ESTS 110C2T7,	Phosphatase
[		GB T42036, AND 110C2XP, GB A1100245; SIMILAR TO	
L			

297	46	MITOCHONDRIAL IMPORT RECEPTOR SUBUNIT Receptor HYPOTHETICAL GB Z98597 FROM S. POMBE. ESTS
		GB T45575 AND GB Z26435 AND GB AA394576 COME
		FROM THIS GENE
301	47	SUBTILISIN PROTEASE STRONG SIMILARITY TO Protease
		PROTEIN SBT1 GB X98929 FROM LYCOPERSICUM
		ESCULENTUM[PUTATIVE]
302	48	ABC TRANSPORTER, MULTI-DRUG RESISTANCE Transporter
		PROTEIN STRONG SIMILARITY TO MRP-LIKE ABC
		TRANSPORTER GB U92650 FROM A. THALIANA AND
		CANALICULAR MULTI-DRUG RESISTANCE PROTEIN
		GB L49379 FROM RATTUS NORVEGICUS
303	49	SERINE/THREONINE PROTEIN PHOSPHATASE Phosphatase
		GB X83099 FROM S. CEREVISIAE[HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
305	50	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE Transferases
		[PUTATIVE]
308		DIMETHYLANILINE MONOOXYGENASE [PUTATIVE] Oxygenases
311	52	ADENYLATE CYCLASE GB AF012921 FROM Cyclase
		MAGNAPORTHE GRISAE. EST GB Z24512 COMES FROM
		THIS GENE; UNKNOWN PROTEIN CONTAINS
210		SIMILARITY TO
312	53	BETA-KETOACYL-COA SYNTHASE STRONG Synthase
		SIMILARITY TO BETA-KETO-COA SYNTHASE
221	EA	GB U37088 FROM SIMMONDSIA CHINENSIS[PUTATIVE]
321	34	ETHYLENE RECEPTOR (ERS2) EST GB W43451 COMES Receptor
324		FROM THIS GENE[PUTATIVE]  1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE Oxidase
324	33	SIMILAR TO ARABIDOPSIS 2A6 (GB X83096). EST
		GB T76913 COMES FROM THIS GENE[PUTATIVE]
327	56	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE Oxidase
321	50	STRONG SIMILARITY TO ARABIDOPSIS 2A6
		(GB X83096)[PUTATIVE]
331	57	ALDO-KETO REDUCTASE BABESIA Reductase
] 551	,	(GB M93122[HYPOTHETICAL PROTEIN SIMILAR TO]

350	58	DIMETHYLANILINE MONOOXYGENASE SIMILAR TO Oxygenases GB:AAC04900[PUTATIVE]
357	59	PECTATE LYASE A11 SIMILAR TO Lyase GB:CAB36835[PUTATIVE]
380	60	SUCROSE-PHOSPHATE SYNTHASE, PUTATIVE SIMILAR Synthase TO GB:Y11795 FROM [CRATEROSTIGMA PLANTAGINEUM]
390	61	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE Oxidase IDENTICAL TO 1-AMINOCYCLOPROPANE-1- CARBOXYLATE OXIDASE (ACC OXIDASE) GB X66719 (EAT1). ESTS GB T43073, GB T5714, GB R90435, GB R44023, GB AA597926, GB A1099676, GB AA650810 AND GB 29725 COME FROM THIS GENE
403	62	O-GLCNAC TRANSFERASE SIMILAR TO RATTUS O-Transferases GLCNAC TRANSFERASE (GB U76557)[PUTATIVE]
412	63	PEROXIDASE ATP12A STRONG SIMILARITY TO Oxidase  ARABIDOPSIS PEROXIDASE ATP11A  (GB X98802)[PUTATIVE]
413	64	PEROXIDASE ATP12A STRONG SIMILARITY TO Oxidase  ARABIDOPSIS PEROXIDASE ATP11A  (GB X98802)[PUTATIVE]
414	65	PEROXIDASE STRONG SIMILARITY TO ARABIDOPSIS Oxidase PEROXIDASE ATPEROX7A (GB X98321)[PUTATIVE]
418	66	ZINC TRANSPORTER SIMILAR TO ARABIDOPSIS FE(II) Transporter TRANSPORT PROTEIN (GB U27590)[PUTATIVE]
419	67	PECTIN METHYLESTERASE SIMILAR TO PRUNUS Esterase PECTINESTERASE (GB X95991)[PUTATIVE]
440	68	UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D-Transferases GLUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D- GLUCOSYLTRANSFERASE GI:2149127 FROM [ARABIDOPSIS THALIANA]
443	69	UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D-Transferases GLUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D- GLUCOSYLTRANSFERASE GI:2149127 FROM [ARABIDOPSIS THALIANA]

444	70	CLUCAN CYNTHACE HIGHLY CIMU AD TO DUTATIVE	Cymthaga
444		GLUCAN SYNTHASE HIGHLY SIMILAR TO PUTATIVE GLUCAN SYNTHASE GB:AAD15408[PUTATIVE]	Symmase
			m
445	71	NA/H ANTIPORTER PROTEINS; N-TERMINAL HALF OF	Transporter
		PROTEIN IS SIMILAR TO NA/H ANTIPORTER	
		PROTEINS[HYPOTHETICAL PROTEIN SIMILAR TO]	
452	72	POLYGALACTURONASE SIMILAR TO	Glycosylase
		GB:AAC23398[PUTATIVE]	
453	73	POLYGALACTURONASE SIMILAR TO	Glycosylase
		GB:AAC23398[PUTATIVE]	
454	74	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		SIMILAR TO INDOLE-3-ACETATE BETA-	
		GLUCOSYLTRANSFERASE GB:AAD32293[PUTATIVE]	
455	75	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		SIMILAR TO INDOLE-3-ACETATE BETA-	
ļ		GLUCOSYLTRANSFERASE GB:AAD32293[PUTATIVE]	
466	76	NPK1-RELATED PROTEIN KINASE 2 PREDICTED BY	Kinase, Protein
		GENEMARK.HMM[PUTATIVE]	
472	77	CHITINASE, CLASS I, PUTATIVE SIMILAR TO	Chitinase
ļ		GB:AAF69774 FROM [ARABIDOPSIS	
ļ		BLEPHAROPHYLLA] (PROC. NATL. ACAD. SCI. U.S.A. 97	
		(10), 5322-5327 (2000))	
486	78	UDPG GLUCOSYLTRANSFERASE GB:AAB62270	Transferases
		GI:2232354 FROM [SOLANUM BERTHAULTII],	
		UNKNOWN PROTEIN CONTAINS SIMILARITY TO	
488	79	FRUCTOKINASE, PUTATIVE SIMILAR TO	Kinase
		FRUCTOKINASE GB:U62329 GI:1915973 FROM	
		[LYCOPERSICON ESCULENTUM]	
489	80		Kinase
		FRUCTOKINASE GB:U37838 GI:1052972 FROM [BETA	-
		VULGARIS]	
494	<b>Q</b> 1	DELTA 9 DESATURASE IDENTICAL TO DELTA 9	Desaturases
774	01	DESATURASE GB:BAA25180 GI:2970034 FROM	
		[ARABIDOPSIS THALIANA]	:
405	00		Decaturação
495	82	DELTA 9 DESATURASE, PUTATIVE SIMILAR TO DELTA	Desaturases
		9 DESATURASE GB:BAA25180 GI:2970034 FROM	
j		[ARABIDOPSIS THALIANA]	

496	83	DELTA 9 DESATURASE, PUTATIVE SIMILAR TO DELTA	Desaturases
		9 DESATURASE GB:BAA25180 GI:2970034 FROM	
		[ARABIDOPSIS THALIANA]	
498	84	DELTA 9 DESATURASE, PUTATIVE SIMILAR TO DELTA	Desaturases
•		9 DESATURASE GB:BAA25180 GI:2970034 FROM	
		[ARABIDOPSIS THALIANA]	
511	85	LIPASE-LIKE PROTEIN SIMILAR TO LIPASE	Lipase
		GB:AAD01804 GI:4103627 FROM [DIANTHUS	
		CARYOPHYLLUS]	
520	86	DELTA 9 DESATURASE, PUTATIVE SIMILAR TO DELTA	Desaturases
		9 DESATURASE GI:2970034 FROM [ARABIDOPSIS	
		THALIANA]	
521	87	DELTA 9 DESATURASE, PUTATIVE SIMILAR TO DELTA	į į
		9 DESATURASE GI:2970034 FROM [ARABIDOPSIS	
		THALIANA]	
532	88	GLUCAN SYNTHASE, PUTATIVE SIMILAR TO GLUCAN	Synthase
		SYNTHASE GB:AAD11794 [FILOBASIDIELLA	
		NEOFORMANS VAR. NEOFORMANS]	
544			Reductase
546			Reductase
547	L		Reductase
552	92	PROTEIN KINASE, 5' PARTIAL CONTAINS PFAM	Kinase, Protein
		PROFILE: PF00069 EUKARYOTIC PROTEIN KINASE	
		DOMAIN	7
555	93	RIBOKINASE, SUGAR KINASE, PUTATIVE SIMILAR TO	Kinase
		RIBOKINASE GB:AAD00536 GI:4099074 FROM	
5.50		[PYROBACULUM AEROPHILUM]	· · · · · · · · · · · · · · · · · · ·
562	94	LIPASE, PUTATIVE SIMILAR TO LIPASE GB:AAD01804	Lipase
		GI:4103627 FROM [DIANTHUS CARYOPHYLLUS]	Dalada
564	95	1	Dehydrogenase
		UNKNOWN PROTEIN CONTAINS PFAM	
		PROFILE:PF02032	7/1
566	96	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		RECEPTOR PROTEIN KINASE GB:BAA11869 GI:1389566	
		FROM [ARABIDOPSIS THALIANA]	

568	97	PROCESSING PEPTIDASE, CHLOROPLAST Protease
		THYLAKOIDAL PROCESSING PEPTIDASE, PUTATIVE
		SIMILAR TO CHLOROPLAST THYLAKOIDAL
		PROCESSING PEPTIDASE GB:CAA71502 GI:2769566
		FROM [ARABIDOPSIS THALIANA]
576	98	NA+/H+ ANTIPORTER GI:2347190 FROM [ARABIDOPSIS Transporter
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
581	99	MITOCHONDRIAL CARRIER Transporter
		PROTEINS[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
596	100	ALTERNATIVE NADH-DEHYDROGENASE GI:3718005 Dehydrogenases
		FROM [YARROWIA LIPOLYTICA] UNKNOWN PROTEIN
		CONTAINS SIMILARITY TO
597	101	PROTEASE ATP-DEPENDENT [HYPOTHETICAL Protease
		PROTEIN CONTAINS SIMILARITY TO]
602	102	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:453245 FROM [MANIHOT ESCULENTA]
603	103	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:453245 FROM [MANIHOT ESCULENTA]
604	104	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:453245 FROM [MANIHOT ESCULENTA]
621	105	PROTEIN PHOSPHATASE 2C, PUTATIVE SIMILAR TO Phosphatase
		GB:CAB90633 FROM [FAGUS SYLVATICA]
622	106	TROPINONE REDUCTASE-I, PUTATIVE SIMILAR TO Reductase
		GB:L20473 FROM [DATURA STRAMONIUM] (PROC.
		NATL. ACAD. SCI. U.S.A. 90, 9591-9595 (1993))
623	107	TROPINONE REDUCTASE-I, PUTATIVE SIMILAR TO Reductase
		GB:L20473 FROM [DATURA STRAMONIUM] (PROC.
		NATL. ACAD. SCI. U.S.A. 90, 9591-9595 (1993))
633	108	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069, EUKARYOTIC PROTEIN KINASE
		DOMAIN (1 COPY)

634	109	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069 EUKARYOTIC PROTEIN KINASE
1		DOMAIN (1 COPY)
642	110	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
042	110	PUTATIVE SIMILAR TO GB:AAC50043 FROM
		[ARABIDOPSIS THALIANA] (PLANT MOL. BIOL. 37 (4),
		<u>-</u> -
(40)	111	587-596 (1998))
649	111	FATTY ACID ELONGASE 3-KETOACYL-COA Synthase
		SYNTHASE, PUTATIVE SIMILAR TO GB:AAC99312
		FROM [ARABIDOPSIS THALIANA] (PLANT J. (1999) IN
		PRESS)
653	112	PHOSPHORIBOSYLANTHRANILATE ISOMERASE Isomerase
		IDENTICAL TO GB:AAB03498 FROM [ARABIDOPSIS
		THALIANA] (CELL 83 (5), 725-734 (1995))
661	113	PROTEIN KINASE, PUTATIVE (FRAGMENT) SIMILAR TO Kinase, Protein
		GB:BAA94509 FROM [POPULUS NIGRA]
663	114	L-ASCORBATE PEROXIDASE IDENTICAL TO Oxidase
		GB:CAA42168 FROM [ARABIDOPSIS THALIANA]
		(PLANT MOL. BIOL. 18 (4), 691-701 (1992))
683	115	HIGH-AFFINITY NITRATE TRANSPORTER NRT2 Transporter
		IDENTICAL TO GB:CAB09794 FROM [ARABIDOPSIS
_		THALIANA]
684	116	HIGH-AFFINITY NITRATE TRANSPORTER ACH2 Transporter
)		IDENTICAL TO GB:AAC35884 FROM [ARABIDOPSIS
		THALIANA] (PLANT J. 17 (5), 563-568 (1999))
695	117	DTDP-GLUCOSE 4-6-DEHYDRATASE GI:9759250 FROM Dehydratase
	!	[ARABIDOPSIS THALIANA] [UNKNOWN PROTEIN
		CONTAINS SIMILARITY TO]
696	118	PROTEASE GI:4415912 FROM [ARABIDOPSIS Protease
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
698	119	AMINO ACID PERMEASE GI:7415521 FROM [ORYZA Transporter
	•	SATIVA][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO
699	120	PREPHENATE DEHYDRATASE GI:1008717 FROM Dehydratase
	120	[AMYCOLATOPSIS METHANOLICA][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
		I KOTEM CONTAINS SIMILARITI TO

702	121	ALPHA2,8-SIALYLTRANSFERASE GI:929684 FROM [MUS MUSCULUS][HYPOTHETICAL PROTEIN CONTAINS	
1 1		SIMILARITY TO]	
724	122	ACYL-(ACYL CARRIER PROTEIN) THIOESTERASE,	Esterase
		PUTATIVE ALMOST IDENTICAL TO ACYL-(ACYL	
		CARRIER PROTEIN) THIOESTERASE GB:CAA85387	
		GI:634003 FROM [ARABIDOPSIS THALIANA]	
725	123	,	Chelatase
}		CHELATASE GB:AF014399 GI:2318116 FROM [PISUM	1
		SATIVUM] VIOLAXANTHIN DE-EPOXIDASE PRECURSOR (U44133)	0.:1
728	124	SIMILAR TO EST GB[N37612[PUTATIVE]	Oxidase
732	125	RECEPTOR KINASE, CLV1 SIMILAR TO RECEPTOR-LIKE	Kinase Protein
/32	123	PROTEIN KINASE (IPOMOEA NIL) (U77888)[PUTATIVE]	Armuse, i lotelli
739	126	BETA-GALACTOSIDE ALPHA-2,3-SIALYLTRANSFERASE	Transferases
		(PIR IJC5251), UNKNOWN PROTEIN SIMILAR TO	
761	127	SUGAR TRANSPORT PROTEIN, ERD6 SIMILAR TO	Transporter
		GB:BAA25989[PUTATIVE]	
769	128	AMIDASE (GB D16207). ESTS	AMIDASE
		GB T20504,GB H36650,GB N97423,GB H36595 COME	,
		FROM THIS GENE; SIMILARITY TO	
772	129	BETA-MANNOSIDASE (GB U46067), UNKNOWN	Glycosylase
		PROTEIN CONTAINS SIMILARITY TO BOS	
773	130	1	Kinase, Protein
		(GB X95577)[HYPOTHETICAL PROTEIN CONTAINS	
785	121	SIMILARITY TO] RATTUS PROTEIN PHOSPHATASE 2C SIMILAR TO	Phosphatase
/63	1.71	GB:AAC16260[PUTATIVE]	i nospiiaiase
792	132		Synthase
	<b>-</b>	GB:BAA74589[PUTATIVE]	
803	133	GALACTINOL SYNTHASE SIMILAR TO GB:AAD26116	Synthase
		FROM [BRASSICA NAPUS][PUTATIVE]	
807	134	LIPASE SIMILAR TO NODULINS AND LIPASE;	Lipase
]		LOCATION OF EST E6C2T7 , GB AA042309. SIMILAR TO	
		NODULINS GI 3328240, GI 2129854 AND OTHERS AND	
		LIPASE, GI 2129636[PUTATIVE]	

- 23 -

900	135	12 OVOBHYTODIENOATE DEDUCTACE ORDI CIVILAD	Poductora
808	135	12-OXOPHYTODIENOATE REDUCTASE OPRI SIMILAR	1
		TO 12-OXOPHYTODIENOATE REDUCTASE, GI 2765083	j
		AND OLD-YELLOW-ENZYME HOMOLOG,	
		GI 2232254[PUTATIVE]	TV:
812	136		Kinase, Protein
		GB:AAD21713[PUTATIVE]	
814	137	BETA-1,3-GLUCANASE GB:AAD22663, LOCATION OF	Glycosylase
		EST 192N12T7, GB R90355, UNKNOWN PROTEIN	
		SIMILAR TO	
823	138	PECTINACETYLESTERASE PRECURSOR SIMILAR TO	
		VIGNA RADIATA PECTINACETYLESTERASE	
		PRECURSOR, GI 1431629[PUTATIVE]	
847	139		Synthase
		GB:CAA07251[PUTATIVE]	
862	140	GLUTAMYL-TRNA REDUCTASE 2 PRECURSOR SIMILAR	Reductase
		TO GB:P49294 AND TO A. THALIANA HEMA2	
		(GB U27118)[PUTATIVE]	
864	141	SUCROSE/H+ SYMPORTER SIMILAR TO GB:CAA76367	Transporter
		AND VICIA SUCROSE TRANSPORT PROTEIN	
		(GB Z93774)[PUTATIVE]	
865	142	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
868	143	REVERSE TRANSCRIPTASE SIMILAR TO	Transcriptase
		GB:AAD29058[PUTATIVE]	
869	144	AMINO ACID PERMEASE GC SPLICE SITE AT POSITION	Transporter
		1256 IS PREDICTED FROM ALIGNMENT AND NOT	
		CONFIRMED EXPERIMENTALLY. HIGHLY SIMILAR TO	
		ARABIDOPSIS THALIANA AMINO ACID PERMEASE I	
		GI 404019, AND OTHER AMINO ACID	
	:	PERMEASES[PUTATIVE]	j
873	145	XYLAN ENDOHYDROLASE SIMILAR TO GB:AAD27896	Hydrolase
		TO ENDOXYLANASES GI 1255238	
	!	(THERMOANAEROBACTERIUM	
		THERMOSULFURIGENES), GI 1813595 (HORDEUM	
	i	VULGARE) AND OTHERS[PUTATIVE]	{
894	146	LECTIN RECEPTOR KINASE VERY SIMILAR TO	Kinase, Protein
	,	GB:CAA69271[PUTATIVE]	
<u></u>		1	

899	147	OXIDOREDUCTASE IN MRPL44-MTF1 INTERGENIC Reductase
		PROTEIN (SP Q05016 YM71_YEAST [UNKNOWN
		PROTEIN SIMILAR TO DAUNORUBICIN C-13 (U77891);
		SIMILAR TO]
904	148	GLUTATHIONE S-TRANSFERASE TSI-1 SIMILAR TO Transferases
	•	GLUTATHIONE S-TRANSFERASE
		(SP Q03666 GTX4_TOBAC); SIMILAR TO EST
		GB H36275[PUTATIVE]
905	149	GLUTATHIONE S-TRANSFERASE TSI-1 SIMILAR TO Transferases
}		GLUTATHIONE S-TRANSFERASE TSI-1 (GI 2190992);
}		SIMILAR TO ESTS GB R29860, EMB Z29757, AND
		EMB Z29758[PUTATIVE]
907	150	CELL RECEPTOR BETA CHAIN CDR3 (GI 3064031); Receptor
		SIMILAR TO NUCLEOPORIN NUP145
		(SP P49687 N145_YEAST); SIMILAR TO ESTS GB N37877,
		EMB Z29159, AND EMB Z30865
908	151	GLUCOSYL TRANSFERASE SIMILAR TO IMMEDIATE-Transferases
		EARLY SALICYLATE-INDUCED  CLUCOSVITE ANISEER ASS (A COOST OF ) PUT ATIVE
913	150	GLUCOSYLTRANSFERASE (AC005167), PUTATIVE  GERMIN-LIKE OXALATE OXIDASE SIMILAR TO ESTS Oxidase
913	132	GB T88481 AND GB AI099566
922	153	ENDOXYLOGLUCAN TRANSFERASE SIMILAR TO Transferases
	100	XYLOGLUCAN ENDOTRANSGLYCOSYLASE-RELATED
		PROTEIN XTR4 (PIR IS71223)[PUTATIVE]
924	154	TYROSINE PHOSPHATASE 2 SIMILAR TO PROTEIN-Phosphatase
		TYROSINE PHOSPHATASE 2 (GI)3249071), SIMILAR TO
		EST GB N96456[PUTATIVE]
929	155	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN Kinase, Protein
		KINASE 1 GI:7573596 FROM [POPULUS NIGRA]
931	156	POLYGALACTURONASE PG1, PUTATIVE SIMILAR TO Glycosylase
		POLYGALACTURONASE PG1 GI:5669846 FROM
		[GLYCINE MAX]
937	157	PHOSPHORIBOSYL DIPHOSPHATE SYNTHASE STRONG Synthase
		SIMILARITY TO GI 4544471 F23E6.11 FROM
		ARABIDOPSIS THALIANA BAC
		GB AC006580[PUTATIVE]

941	158	LIPASE SIMILAR TO GB X02844 LIPASE PRECURSOR	Lipase
		FROM STAPHYLOCOCCUS HYICUS. ESTS GB AI239406	
		AND GB T76725 COME FROM THIS GENE[PUTATIVE]	
943	159	PYRUVATE PHOSPHATE DIKINASE, PEP/PYRUVATE	Kinase
		BINDING DOMAIN	
948	160	ALDO/KETO REDUCTASE FAMILY - AUXIN-INDUCED	Reductase
		PROTEIN STRONG SIMILARITY TO GB X56267 AUXIN-	
		INDUCED PROTEIN (PCNT115) FROM NICOTIANA	
		TABACUM  00248[PUTATIVE]	
952	161	RECEPTOR KINASE, 3' PARTIAL IDENTICAL TO	Kinase, Protein
		GB:AAB65490	
963	162	ZIP4, A PUTATIVE ZINC TRANSPORTER PER	Transporter
		SUGGESTION BY DR. NATASHA M. GROTZ (PNAS, VOL	
		95., 7220-7224)	
969	163	SER/THR PROTEIN KINASE ISOLOG	Kinase, Protein
973	164	LYSOPHOSPHOLIPASE ISOLOG	Lipase
977	165	LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN	Kinase, Protein
		KINASE, PUTATIVE SIMILAR TO LEUCINE-RICH	
		REPEAT TRANSMEMBRANE PROTEIN KINASE	
		GB:AAC27895 GI:3360291 FROM [ZEA MAYS]	
981	166	SECRETORY CARRIER MEMBRANE PROTEIN,	Transporter
}		PUTATIVE SIMILAR TO SECRETORY CARRIER	
	!	MEMBRANE PROTEIN GB:AAF36686 GI:7109228 FROM	
		[ARABIDOPSIS THALIANA]	
982	167	BIFUNCTIONAL NUCLEASE BFN1 ALMOST IDENTICAL	Nuclease
		TO BIFUNCTIONAL NUCLEASE BFN1 GB:AAD00693	
		GI:4099831 FROM [ARABIDOPSIS THALIANA]	
992	168	SERINE/THREONINE KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		SERINE/THREONINE KINASE GB:Y12530 GI:2181187	
		FROM [BRASSICA OLERACEA]	j
994	169	SERINE/THREONINE KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		SERINE/THREONINE KINASE GB:Y12531 GI:2181189	ļ
		FROM [BRASSICA OLERACEA]	
997	170	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR	Kinase, Protein
		TO RECEPTOR-LIKE PROTEIN KINASE GB:AAC95353	1
		GI:4008010 FROM [ARABIDOPSIS THALIANA]	

KINASE 1 GB:BAA23676 GI:2662048 FROM [BRASSICA RAPA]  1001 172 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1279597 FROM [NICOTIANA PLUMBAGINIFOLIA]  1005 173 RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO ESTERASE FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO GBIX95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO GRIAGA04947 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1041 180 PREPHENATE DEHYDRATASE CHLOROPLAST GBIZ1562 AND GBIZ1562 COME FROM THIS GENE[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST GBIZ1562 AND GBIZ1562 COME FROM THIS GENE[PUTATIVE]	998	171	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
1001 172 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO PECTIN METHYLESTERASE GI:1279597 FROM [NICOTIANA PLUMBAGINIFOLIA]  1005 173 RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO KINASE, POTEIN RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA OLERACEA]  1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 KINASE, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO OXIDASE GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY, ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM		ĺ	KINASE 1 GB:BAA23676 GI:2662048 FROM [BRASSICA	
PECTIN METHYLESTERASE GI:1279597 FROM [NICOTIANA PLUMBAGINIFOLIA]  1005 173 RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO KINASE, Protein RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA OLERACEA]  1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO ESTERASE FEUTIN METHYLESTERASE, PUTATIVE SIMILAR TO ESTERASE FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO ESTERASE FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GI/1017583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase (SIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO OXIDASE GI/1617583 FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI/2392772 AND IS A MEMBER OF THE PF[00800 PREPHENATE DEHYDRATASE FAMILY, ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-GI/ycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			RAPA]	
[NICOTIANA PLUMBAGINIFOLIA]  1005 173 RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA OLERACEA]  1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-GI/cosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1001	172	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
1005 173 RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA OLERACEA]  1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-GI)cosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			PECTIN METHYLESTERASE GI:1279597 FROM	
RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA OLERACEA]  1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO G  2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			[NICOTIANA PLUMBAGINIFOLIA]	
OLERACEA]  1008  174  SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019  175  ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022  176  PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023  177  PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035  178  GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040  179  RETICULINE OXIDASE STRONG SIMILARITY TO OXIDASE GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042  180  PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046  181  BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1005	173	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
1008 174 SERINE/THREONINE PROTEIN KINASE EMB CAA69216 Kinase, Protein CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO] 1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM] 1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM] 1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE] 1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE] 1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE] 1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- GB YCOSYIASE BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA	1
CONTAINS SIMILARITY TO EXTENSIN PROTEIN GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GBIX95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GBIAF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			OLERACEA]	:
GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]  1019 175 ENDO-XYLOGLUCAN TRANSFERASE, PUTATIVE Transferases SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GBIX95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GBIAF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI[2392772 AND IS A MEMBER OF THE PF[00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1008	174	SERINE/THREONINE PROTEIN KINASE EMB CAA69216	Kinase, Protein
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SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE GI:2244732 FROM [GOSSYPIUM HIRSUTUM]  1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GI:512380 FROM [BRASSICA NAPUS][SIMILAR TO]	!
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1022 176 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			SIMILAR TO ENDO-XYLOGLUCAN TRANSFERASE	
PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GI:2244732 FROM [GOSSYPIUM HIRSUTUM]	
[LYCOPERSICON ESCULENTUM]  1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO Esterase FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1022	176	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
1023 177 PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			PECTIN METHYLESTERASE GI:1617583 FROM	
FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583 FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			[LYCOPERSICON ESCULENTUM]	
FROM [LYCOPERSICON ESCULENTUM]  1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1023	177	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
1035 178 GLYCOGEN SYNTHASE STRONG SIMILARITY TO Synthase  GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase  GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase  SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			FRUIT-SPECIFIC PECTIN METHYLESTERASE GI:1617583	
GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR (SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			FROM [LYCOPERSICON ESCULENTUM]	
(SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]  1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase  GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase  SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1035	178	GLYCOGEN SYNTHASE STRONG SIMILARITY TO	Synthase
1040 179 RETICULINE OXIDASE STRONG SIMILARITY TO Oxidase  GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase  SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GB X95759 SOLUBLE-STARCH-SYNTHASE PRECURSOR	,
GB AF049347 BERBERINE BRIDGE ENZYME FROM BERBERIS STOLONIFERA[PUTATIVE]  1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			(SSIII) FROM SOLANUM TUBEROSUM[PUTATIVE]	
BERBERIS STOLONIFERA[PUTATIVE]  1042  180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase  SIMILAR TO GI 2392772 AND IS A MEMBER OF THE  PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS  GB T21562 AND GB T21062 COME FROM THIS  GENE[PUTATIVE]  1046  181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase  BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1040	179	RETICULINE OXIDASE STRONG SIMILARITY TO	Oxidase
1042 180 PREPHENATE DEHYDRATASE CHLOROPLAST Dehydratase SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GB AF049347 BERBERINE BRIDGE ENZYME FROM	
SIMILAR TO GI 2392772 AND IS A MEMBER OF THE PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			BERBERIS STOLONIFERA[PUTATIVE]	
PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	1042	180	PREPHENATE DEHYDRATASE CHLOROPLAST	Dehydratase
GB T21562 AND GB T21062 COME FROM THIS GENE[PUTATIVE]  1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			SIMILAR TO GI 2392772 AND IS A MEMBER OF THE	
GENE[PUTATIVE]  1046  181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			PF 00800 PREPHENATE DEHYDRATASE FAMILY. ESTS	
1046 181 BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3- Glycosylase BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GB T21562 AND GB T21062 COME FROM THIS	:
BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM			GENE[PUTATIVE]	
	1046	181	BETA-1,3-GLUCANASE SIMILAR TO GLUCAN ENDO-1,3-	Glycosylase
NICOTIANA TABACUM, ESTS GBIZ18185 AND			BETA-D-GLUCOSIDASE PRECURSOR GB Z28697 FROM	
			NICOTIANA TABACUM. ESTS GB Z18185 AND	Į.
GB AA605362 COME FROM THIS GENE[PUTATIVE]			GB AA605362 COME FROM THIS GENE[PUTATIVE]	

1047	182	LACTOYLGLUTATHIONE LYASE-LIKE PROTEIN Lyase
1017	102	SIMILAR TO PROTEIN GB Z74962 FROM BRASSICA
		OLERACEA WHICH IS SIMILAR TO BACTERIAL YRN1
		AND HEAHIO PROTEINS. ESTS GB T21954, GB T04283,
		GB Z37609, GB N37366, GB R90704, GB F15500 AND
		GB F14353 COME FROM THIS GENE
1055	102	PECTATE LYASE-LIKE PROTEIN SIMILAR TO STYLE Lyase
1055	163	DEVELOPMENT-SPECIFIC PROTEIN 9612 PRECURSOR
		GB X55193 AND PECTATE LYASE P59 PRECURSOR
1062	104	GB X15499 FROM LYCOPERSICON ESCULENTUM
1063	184	PYROPHOSPHATE-FRUCTOSE-6-PHOSPHATE 1-Transferases
		PHOSPHOTRANSFERASE SIMILAR TO
		PYROPHOSPHATE-DEPENDENT
		PHOSPHOFUCTOKINASE BETA SUBUNIT GB Z32850
		FROM RICINUS COMMUNIS. ESTS GB N65773,
		GB N64925 AND GB F15232 COME FROM THIS
1054	107	GENE[PUTATIVE]
1064	185	1-AMINO-CYCLOPROPANE-CARBOXYLIC ACID Oxidase
		OXIDASE (ACC OXIDASE) STRONG SIMILARITY TO
		AMINO-CYCLOPROPANE-CARBOXYLIC ACID OXIDASE
		GB L27664 FROM BRASSICA NAPUS. ESTS GB Z48548
11.50	106	AND GB Z48549 COME FROM THIS GENE[PUTATIVE]
1158	186	NITRATE TRANSPORTER, PUTATIVE SIMILAR TO Transporter
		GB:CAB09794 FROM [ARABIDOPSIS THALIANA]
1163	187	BETA-1,4-N-ACETYLGLUCOSAMINYLTRANSFERASE Transferases
		CONTAINS SIMILARITY TO GI 4417304 F15O11.7
		PUTATIVE BETA-1,4-MANNOSYL-GLYCOPROTEIN
		BETA-1,4-N-ACETYLGLUCOSAMINYLTRANSFERASE
		FROM ARABIDOPSIS THALIANA BAC
		GB AC006446[PUTATIVE]
1170	188	CYTOCHROME P450 MONOOXYGENASE STRONG Oxygenases
		SIMILARITY TO GB D78605 CYTOCHROME P450
		MONOOXYGENASE FROM ARABIDOPSIS THALIANA
		AND IS A MEMBER OF THE PF 00067 CYTOCHROME
		P450 FAMILY[PUTATIVE]

1177	189	CYTOCHROME P450 MONOOXYGENASE STRONG	1
		SIMILARITY TO GI 3313615 F21J9.9 FROM ARABIDOPSIS	l
		THALIANA AND IS A MEMBER OF THE PF 00067	
		CYTOCHROME P450 FAMILY[PUTATIVE]	
1187	190	GLYCOSYL TRANSFERASE GI:6862930 FROM	í
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
	· 	SIMILAR TO][PUTATIVE]	
1190	191	ALLENE OXIDE CYCLASE, PUTATIVE SIMILAR TO	Cyclase
<u> </u>		ALLENE OXIDE CYCLASE GI:8977961 FROM	
		[LYCOPERSICON ESCULENTUM]	
1204	192	STEROID SULFOTRANSFERASE, PUTATIVE SIMILAR TO	i l
		STEROID SULFOTRANSFERASE 1 GI:3420004 FROM	
		[BRASSICA NAPUS]	
1220	193	NADPH-CYTOCHROME P450 REDUCTASE GI:10442765	Reductase
		FROM [TRITICUM AESTIVUM][HYPOTHETICAL	
		PROTEIN CONTAINS SIMILARITY TO]	
1241	194	,	Hydrolase
		HYDROLASE GI:2822275 FROM [PSEUDOMONAS	
		PUTIDA], UNKNOWN PROTEIN CONTAINS SIMILARITY	
10.40	105	TO	Changalogo
1242	195	BETA 1,3-GLUCANASE (GLC1) GI:924952 FROM	Glycosylase
		[TRITICUM AESTIVUM][HYPOTHETICAL PROTEIN	
1260	100	CONTAINS SIMILARITY TO]  RIBULOSE-1,5  BISPHOSPHATE	Tronoforoso
1260	190	CARBOXYLASE/OXYGENASE LARGE SUBUNIT N-	Transferases
		METHYLTRANSFERASE HIGHLY SIMILAR TO	
		RIBULOSE-1,5 BISPHOSPHATE CARBOXYLASE/OXYGENASE LARGE SUBUNIT N-	
		METHYLTRANSFERASE, GI 1731475. RARE GC INTRON	
		SPLICE SITE AT 49572 IS INFERRED FROM PROTEIN	
		ALIGNMENT AND IS NOT CONFIRMED	
		EXPERIMENTALLY[PUTATIVE]	,
1263	197	XYLOGLUCAN FUCOSYLTRANSFERASE, PUTATIVE	Transferases
1 1403	177	SIMILAR TO XYLOGLUCAN FUCOSYLTRANSFERASE	
}			
1265	198	GI:5231145 FROM [ARABIDOPSIS THALIANA]  LIPOAMIDE DEHYDROGENASE, PUTATIVE	Dehydrogenases

1268	199	2-OXOACID DEPENDENT DIOXYGENASE, Oxygenases	
		DIOXYGENASE-LIKE PROTEIN VERY SIMILAR TO	
		ADVENTITIOUS ROOTING RELATED OXYGENASE;	
		VERY SIMILAR TO 2-OXOACID DEPENDENT	
		DIOXYGENASE FROM MALUS DOMESTICA, GI 3492806	
1269	200	2-OXOACID DEPENDENT DIOXYGENASE, Oxygenases	
		DIOXYGENASE-LIKE PROTEIN VERY SIMILAR TO	
		ADVENTITIOUS ROOTING RELATED OXYGENASE;	
]		VERY SIMILAR TO 2-OXOACID DEPENDENT	
		DIOXYGENASE FROM MALUS DOMESTICA, GI 3492806	
1274	201	MANDELONITRILE LYASE SIMILAR TO Reductase	
		MANDELONITRILE LYASES GB:P52707, P52706, AND	
	i	O50048; CONTAINS MC OXIDOREDUCTASES PROTEIN	
		MOTIF[PUTATIVE]	
1275	202	MANDELONITRILE LYASE SIMILAR TO Reductase	
		MANDELONITRILE LYASES GB:P52707, P52706, AND	
	}	O50048; CONTAINS MC OXIDOREDUCTASES PROTEIN	
		MOTIF[PUTATIVE]	
1277	203	RIBONUCLEASE CONTAINS RIBONUCLEASE T2 Nuclease	
,		FAMILY HISTIDINE PROTEIN MOTIF[PUTATIVE]	_
1278	204	RIBONUCLEASE CONTAINS RIBONUCLEASE T2 Nuclease	
		FAMILY HISTIDINE PROTEIN MOTIF[PUTATIVE]	_
1281	205	NUCLEOSIDE TRIPHOSPHATASE, 3' PARTIAL SIMILAR Phosphatase	
1201	206	TO GB:AAC32915[PUTATIVE]	_
1284	206	ACID PHOSPHATASE, PUTATIVE SIMILAR TO ACID Phosphatase	
1004		PHOSPHATASE GI:5360721 FROM [LUPINUS ALBUS]	_
1294	207	RECEPTOR-LIKE PROTEIN KINASE GI:2947063 FROM Kinase, Protei	n
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
1005	200	SIMILAR TO][PUTATIVE]	_
1297	208	PECTATE LYASE, PUTATIVE SIMILAR TO PECTATE Lyase	ļ
1000		LYASE GI:2463509 FROM [ZINNIA ELEGANS]	_
1298	209	GLYOXAL OXIDASE GI:399594 FROM Oxidase	
		[PHANEROCHAETE	
		CHRYSOSPORIUM][HYPOTHETICAL PROTEIN	ĺ
		CONTAINS SIMILARITY TO]	

1301	210	DNA POLYMERASE III GAMMA AND TAU Polymerase
		SUBUNITS[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
1309	211	ANIONIC PEROXIDASE, PUTATIVE SIMILAR TO Oxidase
		ANIONIC PEROXIDASE GI:170202 FROM [NICOTIANA]
		SYLVESTRIS]
1310	212	ANIONIC PEROXIDASE, PUTATIVE SIMILAR TO Oxidase
		ANIONIC PEROXIDASE GI:170202 FROM [NICOTIANA
		SYLVESTRIS]
1321	213	NA+/H+ ANTIPORTER, PUTATIVE SIMILAR TO NA+/H+ Transporter
		ANTIPORTER GB:CAA69925 GI:1655702 FROM
		[XENOPUS LAEVIS]
1326	214	PURPLE ACID PHOSPHATASE, PUTATIVE CONTAINS Phosphatase
		PFAM PROFILE: PF02227 PURPLE ACID PHOSPHATASE
1328	215	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE, Transferases
		PUTATIVE SIMILAR TO XYLOGLUCAN ENDO-
		TRANSGLYCOSYLASE GB:CAA63553 GI:1769907
		[ARABIDOPSIS THALIANA], IDENTICAL TO
		ENDOXYLOGLUCAN TRANSFERASE RELATED
		PROTEIN GB:BAA20290, GI:2154611 [ARABIDOPSIS
		THALIANA]
1337	216	ASPARTATE-SEMIALDEHYDE DEHYDROGENASE, Dehydrogenases
		PUTATIVE SIMILAR TO ASPARTATE-SEMIALDEHYDE
		DEHYDROGENASE SP:031219 [LEGIONELLA
		PNEUMOPHILA]
1345	217	PECTINESTERASE GB:X85216 GI:732912 [PHASEOLUS Esterase
		VULGARIS]; UNKNOWN PROTEIN SIMILAR TO
1361	218	GLUTAMINE AMIDOTRANSFERASE CLASS-IHISTONE Transferases
		ACETYLTRANSFERASE HAT B HYPOTHETICAL
		PROTEIN CONTAINS PFAM PROFILE: PF00117
1362	219	GLYCEROL KINASE IDENTICAL TO SP:P34893 FROM Kinase
		[ARABIDOPSIS THALIANA] (J. MOL. BIOL. 251 (4), 533-
		549 (1995))[PUTATIVE]
1370	220	FAD/NADH-BINDING DOMAIN. ESTS GB H76345 AND Reductase
		GB AA651465 COME FROM THIS GENE
1389		CYSTATHIONINE BETA SYNTHASE DOMAIN Synthase
		[HYPOTHETICAL PROTEIN SIMILAR TO]

1409	222	RECEPTOR LECTIN KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR LECTIN KINASE 3 GI:4100060 FROM
		[ARABIDOPSIS THALIANA]
1410	223	OXIDOREDUCTASE GI:6751707 FROM [ARABIDOPSIS Reductase
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
1411	224	GIBBERELLIN 3 BETA-HYDROXYLASE, PUTATIVE Hydroxylase
		SIMILAR TO GIBBERELLIN 3 BETA-HYDROXYLASE
		GI:3982753 FROM [ARABIDOPSIS THALIANA]
1424	225	INORGANIC PYROPHOSPHATASE, PUTATIVE SIMILAR Phosphatase
		TO INORGANIC PYROPHOSPHATASE GI:790478 FROM
		[NICOTIANA TABACUM]
1450	226	CINNAMOYL COA REDUCTASE, PUITATIVE SIMILAR Reductase
		TO CINNAMOYL COA REDUCTASE GI:2058310 FROM
		[EUCALYPTUS GUNNII]
1454	227	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED Channel
		ION CHANNEL PROTEIN, PUTATIVE SIMILAR TO
		CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED
		ION CHANNEL PROTEIN GI:4581207 FROM
		[ARABIDOPSIS THALIANA]
1464	228	WALL-ASSOCIATED KINASE 2 GI:4826399 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1465	229	WALL-ASSOCIATED KINASE 2 GI:4826399 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1466	230	WALL-ASSOCIATED KINASE 4 GI:3355308 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1467	231	SERINE/THREONINE-SPECIFIC PROTEIN KINASE Kinase, Protein
	İ	GI:7270012 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
1468	232	WALL-ASSOCIATED KINASE 4 GI:3355308 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]

1469	233	WALL-ASSOCIATED KINASE 4 GI:3355308 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1470	234	WALL-ASSOCIATED KINASE 4 GI:3355308 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1475	235	PROTEIN PHOSPHATASE 2C SIMILAR TO Phosphatase
		GB:AAC36699[PUTATIVE]
1479	236	WALL-ASSOCIATED KINASE SIMILAR TO GB AJ012423 Kinase, Protein
		WALL-ASSOCIATED KINASE 2 FROM ARABIDOPSIS
		THALIANA[PUTATIVE]
1487	237	3-DEOXY-D-MANNO-2-OCTULOSONATE-8-PHOSPHATE Synthase
		SYNTHASE SIMILAR TO GB Y14272 3-DEOXY-D-
		MANNO-2-OCTULOSONATE-8-PHOSPHATE SYNTHASE
		FROM PISUM SATIVUM[PUTATIVE]
1491	238	NA/H ANTIPORTER SIMILAR TO GI 4835769 T8K14.18 Transporter
		PUTATIVE NA/H ANTIPORTER ISOLOG FROM
		ARABIDOPSIS THALIANA BAC
1514	220	GB AC007202[PUTATIVE]
1514	239	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein PUTATIVE SIMILAR TO RECEPTOR-LIKE
		PUTATIVE SIMILAR TO RECEPTOR-LIKE SERINE/THREONINE KINASE GI:2465923 FROM
		[ARABIDOPSIS THALIANA]
1525	240	INORGANIC PYROPHOSPHATASE, PUTATIVE SIMILAR Phosphatase
1525	210	TO VACUOLAR-TYPE H+-TRANSLOCATING
		INORGANIC PYROPHOSPHATASE GI:6901678 FROM
		[ARABIDOPSIS THALIANA]
1529	241	H+-ATPASE CATALYTIC SUBUNIT, PUTATIVE SIMILAR ATPase
		TO H+-ATPASE CATALYTIC SUBUNIT GI:6518112 FROM
		[CITRUS UNSHIU]
1546	242	SRG1-LIKE PROTEIN STRONG HOMOLOGY TO SRG1 Oxidase
		PROTEIN, A NEW MEMBER OF THE FE(II)/ASCORBATE
		OXIDASE SUPERFAMILY, 73% IDENTICAL TO SRGI
] ]		[ARABIDOPSIS THALIANA] (GI 479047). LOCATION OF
		ESTS 147E17T7 (GB T76176) AND 136D2T7 (GB T45959)

1547	243	SRG1-LIKE PROTEIN STRONG HOMOLOGY TO SRG1	Oxidase
		PROTEIN, A NEW MEMBER OF THE FE(II)/ASCORBATE	
		OXIDASE SUPERFAMILY, SIMILAR TO SRG1 PROTEIN	
		[ARABIDOPSIS THALIANA] (GI 629561). LOCATION OF	
}		EST F1A5T7 (GB N96370)	
1560	244	POLYGALACTURONASE AFTER FIRST 29 AMINO	Glycosylase
		ACIDS, 43% IDENTICAL TO POLYGALACTURONASE	
		[MEDICAGO SATIVA] (GI 3413322)[PUTATIVE]	
1561	245	RIBOKINASE [PUTATIVE]	Kinase
1562	246	GLUTATHIONE TRANSFERASE ONE OF THREE	Transferases
		REPEATED PUTATIVE GLUTATHIONE TRANSFERASES.	
		72% IDENTICAL TO GLUTATHIONE TRANSFERASE	
		[ARABIDOPSIS THALIANA] (GI 4006934)[PUTATIVE]	
1563	247	GLUTATHIONE TRANSFERASE SECOND OF THREE	Transferases
		REPEATED PUTATIVE GLUTATHIONE TRANSFERASES.	
		72% IDENTICAL TO GLUTATHIONE TRANSFERASE	
		[ARABIDOPSIS THALIANA] (GI 4006934). LOCATION OF	
		ESTS 191A10T7 (GB R90188) AND 171N13T7	
		(GB R65532)[PUTATIVE]	
1564	248	]	Transferases
		REPEATED GLUTATHIONE TRANSFERASES. 65%	
		IDENTICAL TO GLUTATHIONE TRANSFERASE	
		[ARABIDOPSIS THALIANA] (GI 4006934). LOCATION OF	
1568	240	EST 141C5T7 (GB T46669)[PUTATIVE]  LEUCINE-RICH RECEPTOR PROTEIN KINASE 34%	Viscos Dustain
1300	249	IDENTICAL TO LEUCINE-RICH RECEPTOR-LIKE	Kinase, Frotein
		PROTEIN KINASE [IPOMOEA NIL] (GI 1684913) AND 35%	
		IDENTICAL TO LEUCINE-RICH RECEPTOR-LIKE	
		PROTEIN KINASE [MALUS DOMESTICA]	Ì
		(GI 3641252)[PUTATIVE]	
1569	250	RECEPTOR PROTEIN KINASE APPROXIMATELY 30%	Kinase, Protein
		IDENTICAL TO DISEASE RESISTANCE GENES	,
		[LYCOPERSICON PIMPINELLIFOLIUM] (GI 1184077 AND	
		GIJ1184075) AND [LYCOPERSICON ESCULENTUM]	
		(GI 3894387 AND GI 3894393)[PUTATIVE]	<b>[</b>
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1570	251	RECEPTOR PROTEIN KINASE APPROXIMATELY 30% Kinase, Protein
		IDENTICAL TO DISEASE RESISTANCE PROTEINS
		[LYCOPERSICON ESCULENTUM] (GI 3894387 AND
		GI 3894393) AND [LYCOPERSICON PIMPINELLIFOLIUM]
		(GI 1184075 AND GI 1184077)[PUTATIVE]
1584	252	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GB:AAD22368 GI:4544460 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
1600	253	PROTEIN PHOSPHATASE 2C, PUTATIVE SIMILAR TO Phosphatase
		PROTEIN PHOSPHATASE 2C GI:3242077 FROM
		[ARABIDOPSIS THALIANA]
1601	254	PROTEIN PHOSPHATASE 2C, PUTATIVE SIMILAR TO Phosphatase
		PROTEIN PHOSPHATASE 2C GI:3242077 FROM
		[ARABIDOPSIS THALIANA]
1618	255	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR Kinase, Protein
		TO RECEPTOR-LIKE PROTEIN KINASE INRPKI
1634	256	GI:1684913 FROM [IPOMOEA NIL]  WALL-ASSOCIATED KINASE 4 GI:3355308 FROM Kinase, Protein
1034	236	[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1642	257	12-OXOPHYTODIENOATE REDUCTASE, PUTATIVE Reductase
		SIMILAR TO 12-OXOPHYTODIENOATE REDUCTASE
		OPR1 GI:3882355 FROM [ARABIDOPSIS THALIANA]
1644	258	12-OXOPHYTODIENOATE REDUCTASE, PUTATIVE Reductase
		SIMILAR TO 12-OXOPHYTODIENOATE REDUCTASE
		GI:4894182 FROM [LYCOPERSICON ESCULENTUM]
1653	259	THREONYL-TRNA SYNTHETASES [HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
1657	260	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-Isomerase
		TYPE, UNKNOWN PROTEIN CONTAINS PFAM PROFILE:
		PF00254
1669	261	BETA-GLUCAN-ELICITOR RECEPTOR, PUTATIVE Receptor
	!	SIMILAR TO BETA-GLUCAN-ELICITOR RECEPTOR
		GB:D78510 GI:1752733 FROM [GLYCINE MAX]

1674	262	LYSOPHOSPHOLIPASE HOMOLOG, PUTATIVE SIMILAR Lipase
		TO LYSOPHOSPHOLIPASE HOMOLOG GB:AAB97366
		GI:2801536 FROM [ORYZA SATIVA]
1681	263	PEPTIDYL-TRNA HYDROLASE GB:D64003 GI:1001200 Hydrolase
		FROM [SYNECHOCYSTIS SP][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
1683	264	LIPASE GB:AAF36744 GI:7109480 FROM [ARABIDOPSIS Lipase
		THALIANA]; UNKNOWN PROTEIN SIMILAR TO
		PUTATIVE
1687	265	2-ISOPROPYLMALATE SYNTHASE, PUTATIVE SIMILAR Synthase
		TO 2-ISOPROPYLMALATE SYNTHASE GB:AF004165
		GI:2213881 FROM [LYCOPERSICON PENNELLII]
1695	266	GLYCOSYL TRANSFERASE FAMILY 8, HYPOTHETICAL Transferases
		PROTEIN CONTAINS PFAM PROFILE: PF01501
1696	267	FLAVONOL 4'-SULFOTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO FLAVONOL 4'-SULFOTRANSFERASE
		GI:168168 FROM [FLAVERIA CHLORAEFOLIA]
1702	268	BETA-1,3-GLUCANASE-LIKE PROTEIN GI:9758115 FROM Glycosylase
		[ARABIDOPSIS THALIANA], UNKNOWN PROTEIN
		SIMILAR TO
1705	269	REVERSE TRANSCRIPTASE DNA DEPENDENT Transcriptase
		GI:2920563 FROM [SPRAGUEA LOPHII], UNKNOWN
		PROTEIN CONTAINS SIMILARITY TO
1706	270	ALPHA GALACTOSYLTRANSFERASE, PUTATIVE Transferases
	i	SIMILAR TO ALPHA GALACTOSYLTRANSFERASE
		GI:5702018 FROM [TRIGONELLA FOENUM-GRAECUM]
1723	271	ISOCHORISMATE SYNTHASE, PUTATIVE SIMILAR TO Synthase
		ISOCHORISMATE SYNTHASE GI:3348077 FROM
	······································	[ARABIDOPSIS THALIANA]
1739	272	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GI:4006833 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
1743	273	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GI:3738337 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]

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1745	274	[	, Kinase, Protein
Í		PUTATIVE SIMILAR TO RECEPTOR-LIKE	E
ļ		SERINE/THREONINE KINASE GI:2465925 FROM	
		[ARABIDOPSIS THALIANA]	
1753	275	POLYGALACTURONASE-LIKE PROTEIN GI:1017737	Glycosylase
	,	FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	4
		PROTEIN SIMILAR TO]	
1774	276	WALL-ASSOCIATED KINASE 2, PUTATIVE SIMILAR TO	Kinase, Protein
		GB:CAB42872 FROM [ARABIDOPSIS THALIANA]	
		(PLANT MOL. BIOL. 39 (6), 1189-1196 (1999))	
1788	277	2-HYDROXYISOFLAVONE REDUCTASE, PUTATIVE	Reductase
		SIMILAR TO PIR:T08106 FROM [BETULA PENDULA]	
1789	278	GSH-DEPENDENT DEHYDROASCORBATE REDUCTASE	Reductase
		1, PUTATIVE SIMILAR TO GB:BAA90672 FROM [ORYZA	
		SATIVA]	
1790	279	GSH-DEPENDENT DEHYDROASCORBATE REDUCTASE	Reductase
		1, PUTATIVE SIMILAR TO GB:BAA90672 FROM [ORYZA	
		SATIVA]	
1793	280	RIBOKINASE, PUTATIVE SIMILAR TO GB:AAD00536	Kinase
		FROM [PYROBACULUM AEROPHILUM]	
1808	281	PROTEASE LA (LON) ATP-DEPENDENT DOMAIN	Protease
1812	282	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL PROTEIN, PUTATIVE SIMILAR TO	
		CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	
		ION CHANNEL GI:4581207 FROM [ARABIDOPSIS	
		THALIANA]	
1836	283	FERREDOXINNADP REDUCTASE PRECURSOR,	Reductase
		PUTATIVE SIMILAR TO GB:M25528 FROM	
		[MESEMBRYANTHEMUM CRYSTALLINUM]	
1838	284	C-8,7 STEROL ISOMERASE IDENTICAL TO GB:AAD03489	Isomerase
		FROM [ARABIDOPSIS THALIANA] (PLANT MOL. BIOL.	
		38 (5), 807-815 (1998))	
1848	285	SUBTILISIN PROTEASE	Protease
1849	286	SUBTILISIN PROTEASE	Protease
L		<u> </u>	

1863	207	S-ADENOSYL-METHIONINE-STEROL-C-	Transferases
1803	28/		
		METHYLTRANSFERASE IDENTICAL TO STEROL-C-	ļ
		METHYLTRANSFERASE GI:1061040 FROM	
		[ARABIDOPSIS THALIANA]	
1894	288	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE GI:2852447 FROM [ARABIDOPSIS THALIANA]	
1911	289	SUGAR TRANSPORTER PROTEIN NEARLY IDENTICAL	1 -
•		TO ARABIDOPSIS SUGAR TRANSPORTER,	
		GI 1495273[PUTATIVE]	
1913	290	INORGANIC PHOSPHATE TRANSPORTER PROTEIN	Transporter
		SIMILAR TO GB:CAA67395[PUTATIVE]	
1922	291	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE-	Kinase
		DEPENDENT PHOSPHOFRUCTOKINASE ALPHA	
	ĺ	SUBUNIT SIMILAR TO GB:AAC67587 FROM [CITRUS X	
		PARADISI] AND GB:Q41140 FROM [RICINUS	
		COMMUNIS][PUTATIVE]	
1937	292	O-METHYLTRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		GB:AAF28353 FROM [FRAGARIA X ANANASSA]	
1938	293	O-METHYLTRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		GB:AAF28353 FROM [FRAGARIA X ANANASSA]	
1939	294	ATPASE SIMILAR TO GB:AAF28353 FROM [FRAGARIA X	ATPase
		ANANASSA][PUTATIVE]	
1940	295	O-METHYLTRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		GB:AAF28353 FROM [FRAGARIA X ANANASSA]	
1959	296	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
•		GB:AAC63678 GI:3738337 FROM [ARABIDOPSIS	·
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
1968	297	FLAVIN-CONTAINING MONOOXYGENASES, PUTATIVE	Oxygenases
		IDENTICAL TO PUTATIVE FLAVIN-CONTAINING	
		MONOOXYGENASES GB:AAF87896 GI:9454573 FROM	
		[ARABIDOPSIS THALIANA]	
1969	298	CARBOXYVINYL-CARBOXYPHOSPHONATE	Mutase
		PHOSPHORYLMUTASE GB:049290 FROM	
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		SIMILAR TO]	

1988	299	SECA-TYPE CHLOROPLAST PROTEIN TRANSPORT Transporter
		FACTOR SIMILAR TO GB X82404 CHLOROPLAST SECA
		PROTEIN FROM PISUM SATIVUM [PUTATIVE]
2007	300	PECTINESTERASE, PUTATIVE SIMILAR TO Esterase
		PECTINESTERASE GB:CAB08077 GI:1944575 FROM
		[LYCOPERSICON ESCULENTUM]
2008	301	PECTINESTERASE, PUTATIVE SIMILAR TO Esterase
		PECTINESTERASE GB:CAB08077 GI:1944575 FROM
		[LYCOPERSICON ESCULENTUM]
2014	302	PHOSPHATIDYLINOSITOL-4-PHOSPHATE 5-KINASE, Kinase
		PUTATIVE SIMILAR TO PHOSPHATIDYLINOSITOL-4-
		PHOSPHATE 5-KINASE GB:CAB53377 GI:5777366 FROM
		[ARABIDOPSIS THALIANA]
2044	303	TREHALOSE-6-PHOSPHATE PHOSPHATASE, PUTATIVE Phosphatase
		SIMILAR TO TREHALOSE-6-PHOSPHATE
		PHOSPHATASE GI:2944180 FROM [ARABIDOPSIS
		THALIANA]
2051	304	PROTEIN PHOSPHATASE TYPE 2C, PUTATIVE SIMILAR Phosphatase
		TO PROTEIN PHOSPHATASE TYPE 2C GI:4336436 FROM
		[LOTUS JAPONICUS]
2056	305	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:3928543 FROM [ARABIDOPSIS THALIANA]
2057	306	UDP-GLUCOSE GLUCOSYLTRANSFERASE IDENTICAL Transferases
		TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:3928543 FROM [ARABIDOPSIS THALIANA]
2058	307	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:3928543 FROM [ARABIDOPSIS THALIANA]
2059	308	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE, Transferases
		5' PARTIAL SIMILAR TO UDP-GLUCOSE
		GLUCOSYLTRANSFERASE GI:3928543 FROM
		[ARABIDOPSIS THALIANA]
2060	309	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
		GI:6561805 FROM [SORGHUM BICOLOR]
		<u> </u>

2079	310	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE,	Transferases
		PUTATIVE SIMILAR TO	<u> </u> 
		PHOSPHORIBOSYLANTHRANILATE TRANSFERASE	
		GI:1396053 FROM [PISUM SATIVUM]	
2084	311	TRNA ADENYLYLTRANSFERASE SIMILAR TO TRNA	Transferases
	i	ADENYLYLTRANSFERASE GB U15930 FROM LUPINUS	
		ALBUS. EST GB AA721797 COMES FROM THIS	
		GENE[PUTATIVE]	
2089	312	SUCROSE TRANSPORT PROTEIN, SUC2 STRONG	Transporter
		SIMILARITY TO GB:S38196 SUCROSE TRANSPORT	
		PROTEIN SUC2 FROM [ARABIDOPSIS	
		THALIANA][PUTATIVE]	
2090	313	WALL-ASSOCIATED KINASE CONTAINS SIMILARITY	Kinase, Protein
		TO SERINE/THREONINE KINASE GB Y12531 FROM	
		BRASSICA OLERACEA[PUTATIVE]	
2111	314	PHOSPHOMETHYLPYRIMIDINE KINASE PROBABLE	
		THIAMIN BIOSYNTHETIC ENZYME, LOCATION OF EST	
		GB AA395737, GB T21651	
2119	315	NADPH OXIDASE FLAVOCYTOCHROME SUPEROXIDE-	
		GENERATING HIGHLY SIMILAR TO GB:CAA70769,	
		FRO1 AND GB:CAA70770, FRO2 FROM [ARABIDOPSIS	
2125	216	THALIANA][PUTATIVE]	Transporter
2125	310	AUXIN TRANSPORT PROTEIN STRONGLY SIMILAR TO AUXIN TRANSPORT PROTEIN	Transporter
		AUXIN TRANSPORT PROTEIN [GB:AAD52697[PUTATIVE]	
2137	217	PECTINESTERASE SIMILAR TO GB:AAB57669,	Esterace
215/	317	LOCATION OF EST GB Z35063 AND	
		GB Z35062[PUTATIVE]	
2154	318	2-HEXAPRENYL-1,4-NAPHTHOQUINONE	Transferases
		METHYLTRANSFERASE GB:BAA25267 GI:2982680 FROM	
		[MICROCOCCUS LUTEUS]SERINE O-	
	! !	ACETYLTRANSFERASE (EC 2.3.1.30) SAT-52	
		(PIR  S71207) SPORE GERMINATION PROTEIN C2	
		SIMILAR TO	
2164	319	POLYGALACTURONASE, PUTATIVE SIMILAR TO	Glycosylase
		POLYGALACTURONASE GB:BAA88472 GI:6624205	
1		FROM [CUCUMIS SATIVUS]	
L	<u></u>	1	

2165	320	POLYGALACTURONASE PRECURSOR [PUTATIVE]	Glycosylase
2166	321	GLUCOSYLTRANSFERASE GB:AAD15455 GI:4263795	Transferases
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO][PUTATIVE]	
2187	322	CARBONIC ANHYDRASE [PUTATIVE]	Anhydrase
2212	323	REVERSE TRANSCRIPTASE, PUTATIVE SIMILAR TO	Transcriptase
		REVERSE TRANSCRIPTASE GB:BAA20419 GI:2193870	
		FROM [MUS MUSCULUS]	
2220	324	GLUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO	
		BETA-(1-3)-GLUCOSYL TRANSFERASE GB:AAC62210	
	'	GI:3687658 FROM [BRADYRHIZOBIUM JAPONICUM	
2222	325	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		SIMILAR TO GB:AAB64022[PUTATIVE]	-
2223	326	PEROXIDASE SIMILAR TO PEROXIDASE ATP26A,	Oxidase
		GB:CAA72487[PUTATIVE]	
2229	327		Transferases
		GB:AAC78704[PUTATIVE]	
2246	328	i .	Hydroxylase
		BACTERIAL SPECIES[HYPOTHETICAL PROTEIN	
22.50		SIMILAR TO]	
2252	329	LYSINE/HISTIDINE-SPECIFIC PERMEASE SIMILAR TO	Transporter
2252	220	GB:AAC49885, SIMILAR TO EST GB T13994[PUTATIVE]	<u> </u>
2253	330	DEACETYLVINDOLINE 4-O-ACETYLTRANSFERASE,	Transferases
2254	221	GB:AAC9931; UNKNOWN PROTEIN SIMILAR TO  DEACETYLVINDOLINE 4-O-ACETYLTRANSFERASE,	Tronsferen
2234	331	GB:AAC99311[HYPOTHETICAL PROTEIN SIMILAR TO]	Transferases
2256	332	RIBONUCLEASE III SIMILAR TO ESTS EMB Z18464 AND	Nucleos
2230	332	GB AA389811[PUTATIVE]	Nuclease
2271	333	PROTEIN KINASE SIMILAR TO GB:AAD21776, SIMILAR	Kinase Protein
		TO ESTS EMB Z18436, GB T21564, EMB F14127, AND	1211102C, 1 1UtUIII
		GB T75836, DBJ D22341[PUTATIVE]	
2273	334	ANTHRANILATE SYNTHASE BETA SUBUNIT	Synthase
22.5		IDENTICAL TO ANTHRANILATE SYNTHASE BETA	
		SUBUNIT GI:403434 FROM [ARABIDOPSIS THALIANA]	

2277	335	ANTHRANILATE SYNTHASE BETA SUBUNIT Synthase
		IDENTICAL TO ANTHRANILATE SYNTHASE BETA
	٠	SUBUNIT GI:403434 FROM [ARABIDOPSIS THALIANA]
2278	336	PURPLE ACID PHOSPHATASE PRECURSOR GI:7331195 Phosphatase
		FROM [GLYCINE MAX][HYPOTHETICAL PROTEIN
		SIMILAR TO][PUTATIVE]
2294	337	WALL-ASSOCIATED KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		WALL-ASSOCIATED KINASE 1 GI:3549626 FROM
		[ARABIDOPSIS THALIANA]
2296	338	TRNA ISOPENTENYL TRANSFERASE, PUTATIVE Transferases
		SIMILAR TO TRNA ISOPENTENYL TRANSFERASE
		GI:6006718 FROM [ARABIDOPSIS THALIANA]
2301	339	DIHYDROFLAVONOL 4-REDUCTASE, PUTATIVE Reductase
		SIMILAR TO DIHYDROFLAVONOL 4-REDUCTASE
		GI:1332411 FROM [ROSA HYBRIDA]
2308	340	LYSINE AND HISTIDINE SPECIFIC TRANSPORTER, Transporter
		PUTATIVE SIMILAR TO LYSINE AND HISTIDINE
		SPECIFIC TRANSPORTER GI:2576361 FROM
2021		[ARABIDOPSIS THALIANA]
2321	341	ESTERASE 6 GI:606998 FROM [DROSOPHILA Esterase
		SIMULANS], HYPOTHETICAL PROTEIN CONTAINS SIMILAITY TO
2322	242	P-TYPE TRANSPORTING ATPASE, PUTATIVE SIMILAR Transporter
2322	342	TO P-TYPE TRANSPORTING ATPASE, PUTATIVE SIMILAR TRAISPORTER TO P-TYPE TRANSPORTING ATPASE GI:9229867 FROM
		[ARABIDOPSIS THALIANA]
2328	343	URIDINE KINASE GI:6899310 FROM [UREAPLASMA Kinase
2520		UREALYTICUM][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO
2345	344	ALPHA/BETA HYDROLASE GB:AAF67777 GI:7705098 Hydrolase
	•	FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL
		PROTEIN SIMILAR TO][PUTATIVE]
2347	345	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45850 Oxidase
		GI:5262224 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]
2348	346	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45850 Oxidase
		GI:5262224 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]
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2349	347	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45849	Oxidase
		GI:5262223 FROM [ARABIDOPSIS	
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]	
2350	348	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45850	Oxidase
	ė	GI:5262224 FROM [ARABIDOPSIS THALIANA]	
}		UNKNOWN PROTEIN SIMILAR TO	
2351	349	RETICULINE OXIDASE-LIKE PROTEIN GB:CAB45850	Oxidase
		GI:5262224 FROM [ARABIDOPSIS	
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]	
2352	350	FATTY ACID ELONGASE 3-KETOACYL-COA SYNTHASE S	Synthase
		I SIMILAR TO PUTATIVE INTEGRAL MEMBRANE	1
		PROTEIN GB:AAD17424 GI:4335747 FROM	
		[ARABIDOPSIS THALIANA]	
2353	351	BETA-1,3-GLUCANASE GB:AAD26909 GI:4662638 FROM	Glycosylase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		SIMILAR TO][PUTATIVE]	
2389	352	RIBONUCLEASE 3 PRECURSOR IDENTICAL TO	Vuclease
		SP:P42815 FROM [ARABIDOPSIS THALIANA]	
2393	353	REVERSE TRANSCRIPTASE T	ranscriptase
		GB:AAD17398[HYPOTHETICAL PROTEIN SIMILAR TO]	
2402	354	REVERSE TRANSCRIPTASE T	Transcriptase
		GB:AAD22368[HYPOTHETICAL PROTEIN SIMILAR TO]	
2409	355	NITRATE TRANSPORTER, PUTATIVE NITRATE	ransporter
1		TRANSPORTER NTL1 GB:AAC28086 GI:3377517 FROM	
		[ARABIDOPSIS THALIANA]	
2417	356	GALACTOSYLTRANSFERASE, UNKNOWN PROTEIN I	ransferases
		CONTAINS PFAM PROFILE:PF01762	·
2418	357	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO	ransferases
		GLUTATHIONE S-TRANSFERASE GB: AAF22517	
		GI:6652870 FROM [PAPAVER SOMNIFERUM]	
2419	358	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO	ransferases
		GLUTATHIONE S-TRANSFERASE GB: AAF22517	
		GI:6652870 FROM [PAPAVER SOMNIFERUM]	
2421	359	VALYL- TRNA SYNTHETASE, PUTATIVE SIMILAR TO S	ynthase
		VALYL-TRNA SYNTHETASE GB:P93736 GI:3122914	
		FROM [ARABIDOPSIS THALIANA]	

2424	360	RECEPTOR KINASE, PUTATIVE SIMILAR TO CLV1 Kinase, Protein
		RECEPTOR KINASE GB:AAB58929 GI:2160756 FROM
		[ARABIDOPSIS THALIANA]
2437	361	SENSORY TRANSDUCTION HISTIDINE KINASE Kinase, Protein
		SIMILAR TO GB:AAD21777; SIMILAR TO ESTS
		GB AA712891 AND GB AA042438[PUTATIVE]
2474	362	LIPASE SIMILAR TO HYPOTHETICAL PROTEIN Lipase
		GB:AAF24946 GI:6693020 FROM [ARABIDOPSIS
		THALIANA][PUTATIVE]
2494	363	POLYGALACTURONASE GB:AAC04907 GI:2924778 Glycosylase
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL
		PROTEIN SIMILAR TO][PUTATIVE]
2510	364	IRON/ASCORBATE OXIDOREDUCTASE FAMILY Reductase
2516	365	POLY A POLYMERASE FAMILY Polymerase
		MEMBERS[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
2525	366	STEROID SULFOTRANSFERASE, PUTATIVE SIMILAR TO Transferases
		STEROID SULFOTRANSFERASE 3 GI:3420008 FROM
		[BRASSICA NAPUS]
2529	367	PURINE PERMEASE, PUTATIVE SIMILAR TO PURINE Transporter
		PERMEASE GI:7620007 FROM [ARABIDOPSIS
		THALIANA]
2530	368	PURINE PERMEASE IDENTICAL TO PURINE PERMEASE Transporter
		GI:7620007 FROM [ARABIDOPSIS THALIANA]
2542	369	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
25.45	250	KINASE GI:4105699 FROM [ARABIDOPSIS THALIANA]
2547	370	WALL-ASSOCIATED KINASE 1, PUTATIVE SIMILAR TO Kinase, Protein
		WALL-ASSOCIATED KINASE 1 GI:3549626 FROM
2552	271	[ARABIDOPSIS THALIANA]
2552	3/1	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein KINASE GI:4105699 FROM [ARABIDOPSIS THALIANA]
2565	270	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
2303	312	CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
		LIFASE/ACTLMYDRULASE

2566	373	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2567	374	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2568	375	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2569	376	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2571	377	LIPASE, PUTATIVE SIMILAR TO LIPASE GB:AAA93262 Lipase
		GI:1145627 [ARABIDOPSIS THALIANA]
2572	378	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2574	379	LIPASE IDENTICAL TO LIPASE GB:AAA93262 GI:1145627 Lipase
		[ARABIDOPSIS THALIANA] (FEBS LETT. 377 (3), 475-480
		(1995))
2575	380	ANTHRANILATE N-Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,
		PUTATIVE SIMILAR TO ANTHRANILATE N-
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE
2617	201	GB:Z84384 GI:2239084 [DIANTHUS CARYOPHYLLUS]  BETA-1,3 GLUCANASE, PUTATIVE SIMILAR TO Glycosylase
2617	381	
2620	202	GI:7414433 FROM [PISUM SATIVUM]  PHOSPHORIBOSYLANTHRANILATE ISOMERASE Isomerase
2020	362	IDENTICAL TO GI:619749 FROM [ARABIDOPSIS]
		THALIANA] (PLANT CELL 7 (4), 447-461 (1995))
2644	383	REVERSE TRANSCRIPTASE, PUTATIVE SIMILAR TO Transcriptase
2044	202	GI:976278 FROM [ARABIDOPSIS THALIANA]
2645	38/	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
2043	J04	CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
		ENTIODATO I ENTI DIVOLAGE

2646	385	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
[ [		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
2651	386	RECEPTOR-LIKE SERINE/THREONINE KINASE (RFK1), Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE (RFK1) GI:9972369 FROM
		[ARABIDOPSIS THALIANA]
2652	387	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE GI:2465923 FROM
		[ARABIDOPSIS THALIANA]
2653	388	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR PROTEIN KINASE GI:1389566 FROM
		[ARABIDOPSIS THALIANA]
2654	389	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
		PUTATIVE, 5' PARTIAL SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE GI:2465923 FROM
	•	[ARABIDOPSIS THALIANA]
2660	390	PHENYLALANINE HYDROXYLASE GI:476740 FROM Hydroxylase
		[PSEUDOMONAS AERUGINOSA][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
2663	391	ESTERASE GI:4191785 FROM [ARABIDOPSIS Esterase
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
2686	392	BETA-1,3-GLUCANASE PRECURSOR, PUTATIVE Glycosylase
j		SIMILAR TO GI:4097948 FROM [ORYZA SATIVA] (GENE
		223 (1-2), 311-320 (1998))
2688	393	9-CIS-EPOXYCAROTENOID DIOXYGENASE, PUTATIVE Oxygenases
		SIMILAR TO 9-CIS-EPOXYCAROTENOID DIOXYGENASE
		GI:6715257 FROM [PHASEOLUS VULGARIS]
2689	394	DIADENOSINE 5,5-P1,P4-TETRAPHOSPHATE Hydrolase
	!	HYDROLASE, PUTATIVE SIMILAR TO DIADENOSINE
		5,5-P1,P4-TETRAPHOSPHATE HYDROLASE GI:1888556
		FROM [LUPINUS ANGUSTIFOLIUS]
2705	395	CYCLING-ASSOCIATED KINASE GI:1902912 FROM Kinase, Protein
		[RATTUS NORVEGICUS][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]

2707	200	HYDROLASE GI:7270684 FROM [ARABIDOPSIS	TT-vd1
2/0/	396		1 -
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
2710	397	PECTATE LYASE, PUTATIVE CONTAINS PFAM	Lyase
		PROFILE: PF00544: PECTATE LYASE	
2712	398	LIPASE, PUTATIVE CONTAINS PFAM PROFILE: PF01764:	Lipase
		LIPASE	
2714	399	REVERSE TRANSCRIPTASES, POSSIBLE	Transcriptase
		PSEUDOGENE[HYPOTHETICAL PROTEIN SIMILAR TO],	
		PORTIONS OF LINE-ELEMENT	
2716	400	ABC TRANSPORTER, PUTATIVE CONTAINS PFAM	Transporter
		PROFILE: PF00005: ABC TRANSPORTER	
2717	401	ABC TRANSPORTER, PUTATIVE CONTAINS PFAM	Transporter
į		PROFILE: PF00005: ABC TRANSPORTER	
2725	402	FERRODOXIN NADP OXIDOREDUCTASE, PUTATIVE	Reductase
		SIMILAR TO FERRODOXIN NADP OXIDOREDUCTASE	
		GB:X99419 GI:1480346 FROM [PISUM SATIVUM]	
2727	403	UDP GLUCOSE:FLAVONOID 3-O-	Transferases
		GLUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO UDP	
		GLUCOSE:FLAVONOID 3-O-GLUCOSYLTRANSFERASE	
		GB:AAB81683 GI:2564114 FROM [VITIS VINIFERA]	
2731	404	SERINE/THREONINE PROTEIN KINASE CONTAINS	Kinase, Protein
		EUKARYOTIC PROTEIN KINASE DOMAIN PF 00069,	ŕ
		SIMILAR TO GB:AAB47421[PUTATIVE]	·
2744	405	RETICULINE OXIDASE-LIKE PROTEIN SIMILAR TO	Oxidase
		GB:P30986 FROM [ESCHSCHOLZIA CALIFORNICA] (	
		BERBERINE BRIDGE-FORMING ENZYME ), ESTS	i
		GB F19886, GB Z30784 AND GB Z30785 COME FROM THIS	
		GENE[PUTATIVE]	
2745	406	RETICULINE OXIDASE-LIKE PROTEIN SIMILAR TO	Ovidase
2143	400	GB:P30986 FROM [ESCHSCHOLZIA CALIFORNICA] (	Calduse
			Ì
		GB F19886, GB Z30784 AND GB Z30785 COME FROM THIS	`
		GENE[PUTATIVE]	

2746	407	RETICULINE OXIDASE-LIKE PROTEIN SIMILAR TO	Oxidase
		GB:P30986 FROM [ESCHSCHOLZIA CALIFORNICA] (	
		BERBERINE BRIDGE-FORMING ENZYME ), ESTS	
		GB F19886, GB Z30784 AND GB Z30785 COME FROM THIS	
		GENE[PUTATIVE]	
2747	408	RETICULINE OXIDASE-LIKE PROTEIN SIMILAR TO	Oxidase
		GB:P30986 FROM [ESCHSCHOLZIA CALIFORNICA] (	
		BERBERINE BRIDGE-FORMING ENZYME ), ESTS	ļ
		GB F19886, GB Z30784 AND GB Z30785 COME FROM THIS	
		GENE[PUTATIVE]	
2748	409	RETICULINE OXIDASE-LIKE PROTEIN SIMILAR TO	Oxidase
		GB:P30986 FROM [ESCHSCHOLZIA CALIFORNICA] (	·
		BERBERINE BRIDGE-FORMING ENZYME ), ESTS	
		GB F19886, GB Z30784 AND GB Z30785 COME FROM THIS	
		GENE[PUTATIVE]	
2750	410	RETICULINE OXIDASE-LIKE PROTEIN, 3' PARTIAL	Oxidase
		SIMILAR TO GB:P30986 FROM [ESCHSCHOLZIA]	
		CALIFORNICA] ( BERBERINE BRIDGE-FORMING	
		ENZYME ), ESTS GB F19886, GB Z30784 AND GB Z30785	
		COME FROM THIS GENE[PUTATIVE]	
2759	411	PEROXIDASE SIMILAR TO CATIONIC PEROXIDASE	Oxidase
		(GI 1232069); SIMILAR TO EST GB AI100412[PUTATIVE]	
2762	412	VACUOLAR SORTING RECEPTOR SIMILAR TO	Receptor
		(GI 3033390); SIMILAR TO EST DBJ C72582[PUTATIVE]	
2784	413	POTASSIUM TRANSPORTER SIMILAR TO POTASSIUM	Transporter
2501	41.	TRANSPORTER GB:AAB87687[PUTATIVE]	
2794	414		Transferases
		FORMYLTRANSFERASE ALMOST IDENTICAL TO	
		GB:P52422 FROM [ARABIDOPSIS THALIANA],	
0705	415	INVOLVED WITH PURINE BIOSYNTHESIS[PUTATIVE]	Dobardage
2795	415	ASPARTATE KINASE-HOMOSERINE DEHYDROGENASE	Denydrogenases
		ALMOST IDENTICAL TO GB X71364 GENE FOR	
		ASPARTATE KINASE HOMOSERINE DEHYDROGENASE	
<u> </u>		FROM ARABIDOPSIS THALIANA[PUTATIVE]	

2798	416	IRON-REGULATED TRANSPORTER PROTEIN	Transporter
2/90	410	PUTATIVE SIMILAR TO IRON-REGULATED	1 -
	}	TRANSPORTER 2 GB:AAD30549 GI:4836773 FROM	
		[LYCOPERSICON ESCULENTUM]	
2814	417	PROTEIN KINASE, PUTATIVE CONTAINS PFAM	(
	Ì	PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE	
	·	DOMAIN	
2821	418	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	Transferases
	}	PUTATIVE SIMILAR TO N-	
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE	
	i	[IPOMOEA BATATAS] GI:6469032	
2836	419	COPPER AMINE OXIDASE, PUTATIVE SIMILAR TO	Oxidase
		COPPER AMINE OXIDASE GI:4651202 FROM [PISUM	
1		SATIVUM]	
2838	420	COPPER AMINE OXIDASE, PUTATIVE SIMILAR TO	Oxidase
		COPPER AMINE OXIDASE GI:685197 FROM [PISUM	
		SATIVUM]	
2853	421	AMINO ACID PERMEASE, PUTATIVE CONTAINS PFAM	Transporter
		PROFILE: PF00324: AMINO ACID PERMEASE	
2854	422	AMINO ACID PERMEASE, PUTATIVE CONTAINS PFAM	Transporter
		PROFILE: PF00324: AMINO ACID PERMEASE	
2857	423	PHOSPHORIBOSYL-ATP PYROPHOSPHOHYDROLASE	Hydrolase
		(AT-IE) IDENTICAL TO PHOSPHORIBOSYL-ATP	-
		PYROPHOSPHOHYDROLASE (AT-IE) [ARABIDOPSIS	
		THALIANA] GI:3461884 (PLANT PHYSIOL. 118 (1), 275-	
		283 (1998))	
2859	474	MAJOR INTRINSIC PROTEIN, PUTATIVE CONTAINS	Channel
2039	727	TIGRFAM PROFILE: TIGR00861: MIP FAMILY CHANNEL	
		PROTEINS	·
2860	425	MAJOR INTRINSIC PROTEIN, PUTATIVE CONTAINS	Channel
2000	423	· ·	Chame
		TIGRFAM PROFILE: TIGR00861: MIP FAMILY CHANNEL	
2005	10 -	PROTEINS	G. d.
2865	426	TERPENE SYNTHASE FAMILY PROTEIN, PUTATIVE	Synthase
]		CONTAINS PFAM PROFILE: PF01397: TERPENE	
		SYNTHASE FAMILY	

2873	427	SECRETORY CARRIER MEMBRANE PROTEIN, Transporter
		PUTATIVE SIMILAR TO SECRETORY CARRIER
		MEMBRANE PROTEIN GI:7109228 FROM [ARABIDOPSIS]
		THALIANA]
2874	428	PHOSPHORIBULOKINASE PRECURSOR IDENTICAL TO Kinase
		PHOSPHORIBULOKINASE PRECURSOR GB:P25697
		GI:125576 FROM [ARABIDOPSIS THALIANA]
2878	429	PINORESINOL-LARICIRESINOL REDUCTASE, PUTATIVE Reductase
		SIMILAR TO PINORESINOL-LARICIRESINOL
		REDUCTASE GB:AAF63508 GI:7542583 FROM [THUJA
		PLICATA]
2883	430	OBTUSIFOLIOL 14-ALPHA-DEMETHYLASE (CYP51) Methylase
		GB:Y09292 GI:1707854 FROM [TRITICUM AESTIVUM],
		CONTAINS SIMILARITY TO
2884	431	ENDOXYLOGLUCAN TRANSFERASE, PUTATIVE Transferases
		SIMILAR TO ENDOXYLOGLUCAN TRANSFERASE
		GB:AAD45125 GI:5533313 FROM [ARABIDOPSIS
		THALIANA]
2885	432	CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase
		PUTATIVE SIMILAR TO CELLULOSE SYNTHASE
		CATALYTIC SUBUNIT GB:BAB09693 GI:9759258 FROM
		[ARABIDOPSIS THALIANA]
2887	433	GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE Transferases
		IDENTICAL TO GLYCEROL-3-PHOSPHATE
		ACYLTRANSFERASE GB:Q43307 FROM [ARABIDOPSIS
		THALIANA]
2897	434	L-GULONOLACTONE OXIDASE GB:J03536 GI:204149 Oxidase
		FROM [RATTUS NORVEGICUS] UNKNOWN PROTEIN
		CONTAINS SIMILARITY TO
2898	435	L-GULONOLACTONE OXIDASE GB:J03536 GI:204149 Oxidase
2000	700	FROM [RATTUS NORVEGICUS] UNKNOWN PROTEIN
		CONTAINS SIMILARITY TO
2902	126	OXIDASE, PUTATIVE SIMILAR TO OXIDASE Oxidase
2902	450	
		GB:AAA32870 GI:166876 FROM [ARABIDOPSIS
		THALIANA]

2930	127	PROTEIN KINASE, PUTATIVE IDENTICAL TO BHLH Kinase, Protein
2930	437	
		PROTEIN GB:CAA67885 GI:1465368 FROM
		[ARABIDOPSIS THALIANA]
2938	438	PROTEIN PHOSPHATASE 2C GB:CAA72341 GI:2582800 Phosphatase
	!	FROM [MEDICAGO SATIVA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
2952	439	BETA-1,3-GLUCANASE PRECURSOR, PUTATIVE Glycosylase
		SIMILAR TO BETA-1,3-GLUCANASE PRECURSOR
		GI:4097948 FROM [ORYZA SATIVA]
2955	440	STARCH SYNTHASE, PUTATIVE SIMILAR TO STARCH Synthase
		SYNTHASE GI:21613 FROM [SOLANUM TUBEROSUM]
2965	441	NON-LTR RETROELEMENT REVERSE Transcriptase
		TRANSCRIPTASE[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO][PUTATIVE]
2968	442	CATECHOL O-METHYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO CATECHOL O-METHYLTRANSFERASE
		GI:4808524 FROM [THALICTRUM TUBEROSUM]
2981	443	BETA-1,3-GLUCANASE, PUTATIVE SIMILAR TO ENDO-Glycosylase
2>01		1,3-BETA-GLUCANASE GB:AAC39322 GI:2735502 FROM
		[HORDEUM VULGARE]
2985	111	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
2963	444	PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE
		DOMAIN
2999	445	NITRATE TRANSPORTER NTL1, PUTATIVE SIMILAR TO Transporter
		GI:3377517 FROM [ARABIDOPSIS THALIANA]
3015	446	EPIDERMAL GROWTH FACTOR RECEPTOR-LIKE Receptor
		PROTEIN GB:AAB31972 GI:9256501 FROM
		[XIPHOPHORUS MACULATUS][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
3017	447	DISEASE RESISTANCE PROTEIN, PUTATIVE SIMILAR Kinase, Protein
		TO RECEPTOR KINASE-LIKE PROTEIN GB:AAB82755
		GI:2586083 FROM [ORYZA LONGISTAMINATA]
		(SCIENCE 270 (5243), 1804-1806 (1995))
3024	448	TERPENE SYNTHASE, PUTATIVE SIMILAR TO DELTA-Synthase
		CADINENE SYNTHASE ISOZYME A GB:Q43714 FROM
		[GOSSYPIUM ARBOREUM]

3050	440	POLYNEURIDINE ALDEHYDE ESTERASE, PUTATIVE Esterase
3030	449	·
		SIMILAR TO POLYNEURIDINE ALDEHYDE ESTERASE
		GI:6651393 FROM [RAUVOLFIA SERPENTINA]
3062	450	RECEPTOR PROTEIN KINASE-LIKE PROTEIN Kinase, Protein
		GI:10177178 FROM [ARABIDOPSIS]
		THALIANA][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
3066	451	PSEUDOURIDINE SYNTHASES[PUTATIVE], PROTEIN Synthase
		CONTAINS SIMILARITY TO
3072	452	SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO SOMATIC EMBRYOGENESIS
		RECEPTOR-LIKE KINASE GI:2224910 FROM [DAUCUS
		CAROTA]
3079	453	RECEPTOR-LIKE PROTEIN KINASE GI:6979335 FROM Kinase, Protein
		[ORYZA SATIVA][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
3082	454	PEROXIDASE, PUTATIVE SIMILAR TO GI:1199777 FROM Oxidase
		[POPULUS NIGRA]
3087	455	DNA POLYMERASE I, PUTATIVE SIMILAR TO Polymerase
		GI:4090935 FROM [RHODOTHERMUS SP. 'ITI 518']
3091	456	PROTEIN KINASE DOMAIN (2 COPIES), PF00560 Kinase, Protein
		LEUCINE RICH REPEAT (17 COPIES)
3099	457	PEROXIDASE ATP13A, PUTATIVE SIMILAR TO Oxidase
		GB:CAA67312 FROM [ARABIDOPSIS THALIANA]
3121	458	PROTEIN PHOSPHATASE TYPE 2C, PUTATIVE SIMILAR Phosphatase
		TO GB:AAD17805 FROM [LOTUS JAPONICUS] (PROC.
		NATL. ACAD. SCI. U.S.A. 96 (4), 1738-1743 (1999))
3143	459	HYOSCYAMINE 6-DIOXYGENASE HYDROXYLASE, Oxygenases
		PUTATIVE SIMILAR TO HYOSCYAMINE 6-
		DIOXYGENASE HYDROXYLASE GB:P24397 FROM
		[HYOSCYAMUS NIGER]
3199	460	PROTEIN KINASE, PUTATIVE SIMILAR TO MANY Kinase, Protein
		PREDICTED PROTEIN KINASES
3214	461	TREHALOSE-PHOSPHATASE, PUTATIVE CONTAINS Phosphatase
	-101	TIGRFAM PROFILE: TIGR00685: TREHALOSE-
		PHOSPHATASE
		HOSHMASE

3239	462	ACETYL-COA CARBOXYLASE, PUTATIVE SIMILAR TO Carboxylase
		GI:1100253 FROM [ARABIDOPSIS THALIANA]
3244	463	ADENYLOSUCCINATE LYASE-LIKE PROTEIN SIMILAR Lyase
		TO SP:P44797 FROM [HAEMOPHILUS INFLUENZAE]
3249	464	VANILLOID RECEPTOR-LIKE PROTEIN GB:AAD26363 Receptor
•		FROM [HOMO SAPIENS][HYPOTHETICAL PROTEIN
		SIMILAR TO]
3284	465	NON-LTR REVERSE TRANSCRIPTASES[HYPOTHETICAL Transcriptase
		PROTEIN CONTAINS SIMILARITY TO]
3290	466	BIOTIN HOLOCARBOXYLASE SYNTHETASE, PUTATIVE Synthase
		SIMILAR TO BIOTIN HOLOCARBOXYLASE
		SYNTHETASE GI:4874309 FROM [ARABIDOPSIS
		THALIANA]
3340	467	PECTINESTERASE, PUTATIVE, 5' PARTIAL SIMLAR TO Esterase
		GI:1944574 FROM [LYCOPERSICON ESCULENTUM]
3394	468	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase
		POLYGALACTURONASE GI:288611 FROM [ZEA MAYS]
3395	469	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase
		POLYGALACTURONASE GI:288611 FROM [ZEA MAYS]
3396	470	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase
	-	POLYGALACTURONASE GI:288611 FROM [ZEA MAYS]
3413	471	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GI:4006833 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
3418	470	TO][PUTATIVE]  STEROL GLUCOSYLTRANSFERASE, PUTATIVE Transferases
3418		
	412	
	412	SIMILAR TO STEROL GLUCOSYLTRANSFERASE
3427		SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM]
3427		SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM] HISTIDINE DECARBOXYLASE, PUTATIVE SIMILAR TO Decarboxylase
3427		SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM] HISTIDINE DECARBOXYLASE, PUTATIVE SIMILAR TO Decarboxylase HISTIDINE DECARBOXYLASE GB:BAA78331 GI:4996105
	473	SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM] HISTIDINE DECARBOXYLASE, PUTATIVE SIMILAR TO Decarboxylase HISTIDINE DECARBOXYLASE GB:BAA78331 GI:4996105 FROM [BRASSICA NAPUS]
3427	473	SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM] HISTIDINE DECARBOXYLASE, PUTATIVE SIMILAR TO Decarboxylase HISTIDINE DECARBOXYLASE GB:BAA78331 GI:4996105 FROM [BRASSICA NAPUS] REVERSE TRANSCRIPTASE, PUTATIVE SIMILAR TO Transcriptase
	473 474	SIMILAR TO STEROL GLUCOSYLTRANSFERASE GI:4731867 FROM [DICTYOSTELIUM DISCOIDEUM] HISTIDINE DECARBOXYLASE, PUTATIVE SIMILAR TO Decarboxylase HISTIDINE DECARBOXYLASE GB:BAA78331 GI:4996105 FROM [BRASSICA NAPUS]

3439	475	STEAROYL ACYL CARRIER PROTEIN DESATURASE,	Desetureses
3437	4/5	PUTATIVE SIMILAR TO STEAROYL ACYL CARRIER	
		ł	ł
		PROTEIN DESATURAȘE LLDD3A20 GB: AAD28287	
		GI:4704824 FROM [LUPINUS LUTEUS]	
3455	476	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	-
		GI:4544460 FROM [ARABIDOPSIS THALIANA],	
		UNKNOWN PROTEIN SIMILAR TO PUTATIVE	
3464	477	GIBBERELLIN 20-OXIDASE, PUTATIVE, 5' PARTIAL	Oxidase
		SIMILAR TO GIBBERELIN 20-OXIDASE GI:4164141	
		FROM [LACTUCA SATIVA]	
3470	478	AMINO ACID PERMEASE, PUTATIVE ALMOST	Transporter
		IDENTICAL TO AMINO ACID PERMEASE GI:608673	
		FROM [ARABIDOPSIS THALIANA]	
3471	479	GIBBERELLIN 20-OXIDASE, PUTATIVE SIMILAR TO	Oxidase
		GIBBERELIN 20-OXIDASE GI:4164141 FROM [LACTUCA	
		SATIVA]	
3497	480	PEROXIDASE ATP18A, 3' PARTIAL IDENTICAL TO	Oxidase
		GB:CAA67336 FROM [ARABIDOPSIS THALIANA]	
3498	481	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
		PECTIN METHYLESTERASE GI:1617588 FROM	
		[LYCOPERSICON ESCULENTUM]	
3537	482	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		GI:3746069 FROM [ARABIDOPSIS	-
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TOJ[PUTATIVE]	
3544	483	PHYTOCHELATIN SYNTHETASE GI:3559805 FROM	Synthase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	•
		SIMILAR TO][PUTATIVE]	Ì
3545	484	AAA-TYPE ATPASE-LIKE PROTEIN GI:9759053 FROM	ATPase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		SIMILAR TO	
3564	195	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptoca
3304	402	GB:AAD21515 GI:4510429 FROM [ARABIDOPSIS]	11anscriptase
		`	
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
2570	400	TO][PUTATIVE]	Dh a mh a ta a
3570		PROTEIN PHOSPHATASE 2C GB:AAD25933 GI:4587992	rnospnatase
		FROM [ARABIDOPSIS THALIANA]	

3578	487	ESTERASE GI:4335745 FROM [ARABIDOPSIS	Esterase
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	}
		TO][PUTATIVE]	
2.707	100		
3597	488	LYSINE AND HISTIDINE SPECIFIC TRANSPORTER,	_
		PUTATIVE SIMILAR TO LYSINE AND HISTIDINE	
		SPECIFIC TRANSPORTER GI:2576361 FROM	
		[ARABIDOPSIS THALIANA]	
3610	489	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		GB:AAC63678 GI:3738337 FROM [ARABIDOPSIS	
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
3632	490	DIOXYGENASE, PUTATIVE SIMILAR TO DIOXYGENASE	Oxygenases
		GB:CAA70330 GI:1666096 FROM [MARAH	
		MACROCARPUS]	,
3643	491	POLYGALACTURONASE PG1, PUTATIVE SIMILAR TO	Glycosylase
	-	GB:AAD46483 FROM [GLYCINE MAX] (MOL. PLANT	,
		MICROBE INTERACT. 12 (6), 490-498 (1999))	
3660	402	G PROTEIN COUPLED RECEPTOR IDENTICAL TO	Recentor
3000	772	PUTATIVE G PROTEIN COUPLED RECEPTOR GI:2104224	Receptor
		FROM [ARABIDOPSIS THALIANA][PUTATIVE]	
3674	402		D
3074	493		Deaminase
		DEAMINASE [UNKNOWN PROTEIN CONTAINS	
		SIMILARITY TO]	
3680	494	PROTEIN KINASE, PUTATIVE CONTAINS SIMILARITY	Kinase, Protein
		TO MANY PREDICTED PROTEIN KINASES	
3696		LYSINE AND HISTIDINE SPECIFIC TRANSPORTER,	_
		PUTATIVE SIMILAR TO LYSINE AND HISTIDINE	
		SPECIFIC TRANSPORTER GI:2576361 FROM	
		[ARABIDOPSIS THALIANA]	
3703	496	LECTIN RECEPTOR KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		LECTIN RECEPTOR KINASE GI:1769897 FROM	
		[ARABIDOPSIS THALIANA]	
3711	497	TERPENE CYCLASE, PUTATIVE SIMILAR TO TERPENE	Cyclase
		CYCLASE GI:9293912 FROM [ARABIDOPSIS THALIANA]	
3712	498	TERPENE CYCLASE, PUTATIVE SIMILAR TO TERPENE	Cyclase
		CYCLASE GI:9293912 FROM [ARABIDOPSIS THALIANA]	-
J			

3715	499	CHORISMATE SYNTHASE, PUTATIVE SIMILAR TO Synthase
		CHORISMATE SYNTHASE GI:452796 FROM
		[SYNECHOCYSTIS SP.]
3720	500	FLAVIN-CONTAINING MONOOXYGENASE, PUTATIVE Oxygenases
		SIMILAR TO FLAVIN-CONTAINING MONOOXYGENASE
		4 GI:31429 FROM [HOMO SAPIENS]
3736	501	LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO LIGHT REPRESSIBLE
		RECEPTOR PROTEIN KINASE GI:1321686 FROM
		[ARABIDOPSIS THALIANA]
3753	502	RECEPTOR-LIKE KINASE GI:2224910 FROM [DAUCUS Kinase, Protein
		CAROTA][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
3764	503	FLAVONOL SYNTHASE, PUTATIVE SIMILAR TO Synthase
		FLAVONOL SYNTHASE GI:311657 FROM [PETUNIA]
		HYBRIDA]
3776	504	GERANYL GERANYL PYROPHOSPHATE SYNTHASE, Synthase
		PUTATIVE SIMILAR TO GERANYL GERANYL
		PYROPHOSPHATE SYNTHASE GB:BAA23157 GI:2578822
2790	505	FROM [ARABIDOPSIS THALIANA]
3780	505	PEROXIDASE, PUTATIVE SIMILAR TO PEROXIDASE Oxidase
3787	506	GB:P80679 FROM [ARMORACIA RUSTICANA]  ESTERASE GB:AAD17422 GI:4335745 FROM Esterase
3/6/	300	[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO][PUTATIVE]
3788	507	ESTERASE GB:AAD17422 GI:4335745 FROM Esterase
3768	307	[ARABIDOPSIS THALIANA; UNKNOWN PROTEIN
		SIMILAR TO
3789	508	ESTERASE GB:AAD17422 GI:4335745 FROM Esterase
		[ARABIDOPSIS THALIANA]ESTERASE GB:AAD17422
		GI:4335745 FROM [ARABIDOPSIS THALIANA;
		UNKNOWN PROTEIN SIMILAR TO UNKNOWN PROTEIN
		SIMILAR TO
3793	509	FUCOSYLTRANSFERASE C3 PROTEIN, PUTATIVE Transferases
,		SIMILAR TO FUCOSYLTRANSFERASE C3 PROTEIN
1		
		GB:CAB52254 GI:5702039 FROM [VIGNA RADIATA]

3796	510	MAP3K-LIKE PROTEIN KINASE GB:CAB16796	Kinase, Protein
		GI:4006878 FROM [ARABIDOPSIS THALIANA]	
3801	511	GLUTAMINE AMIDOTRANSFERASES CLASS-	Transferases
		IIGLUTAMINE AMIDOTRANSFERASE CLASS-IHISTONE	
		ACETYLTRANSFERASE HAT B HYPOTHETICAL	
		PROTEIN CONTAINS PFAM PROFILE: PF00117	
		HYPOTHETICAL PROTEIN CONTAINS PFAM PROFILE:	
		PF00310	
3803	512	NA+/H+ ANTIPORTER, PUTATIVE SIMILAR TO PROTEIN	Transporter
		CONTAINED WITHIN GB:AE001273 FROM [CHLAMYDIA	
		TRACHOMATIS]	
3808	513	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR	Transferases
		TO GI:860955 FROM [HYOSCYAMUS MUTICUS] (PLANT	
		PHYSIOL. 109 (1), 253-260 (1995))	
3825			Kinase
	•	GB:AAD16273 GI:4323240 FROM [FILOBASIDIELLA	
		NEOFORMANS], UNKNOWN PROTEIN CONTAINS	
3828	£15	SIMILARITY TO BRANCHED-CHAIN AMINO ACID	Transferases
3828	313	AMINOTRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		BRANCHED-CHAIN AMINO ACID AMINOTRANSFERASE	
		GB:AAF07191 GI:6319165 FROM [SOLANUM]	
		TUBEROSUM]	
3835	516		Transcriptase
		TRANSCRIPTASE GB:AAD25836 GI:4587608 FROM	•
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		SIMILAR TO][PUTATIVE]	
3856	517	FRUCTOKINASE, PUTATIVE SIMILAR TO	Kinase
		FRUCTOKINASE GB:U37838 GI:1052972 FROM [BETA	·
		VULGARIS]	
3873	518	UDP RHAMNOSE: ANTHOCYANIDIN-3-GLUCOSIDE	Transferases
		RHAMNOSYLTRANSFERASE, PUTATIVE SIMILAR TO	
		UDP RHAMNOSE: ANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE [PETUNIA X HYBRIDA]	
		GI:397567	

3876	519	PROTEIN KINASE, PUTATIVE CONTAINS PFAM	Kinase, Protein
		PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE	-
		DOMAIN	
3898	520	DNA POLYMERASE A FAMILY PROTEIN, PUTATIVE	Polymerase
		CONTAINS PFAM PROFILE: PF00476: DNA	
		POLYMERASE FAMILY A	
3909	521	GIBBERELLIN 20-OXIDASE, PUTATIVE SIMILAR TO	Oxidase
		GIBBERELLIN 20-OXIDASE [TRITICUM AESTIVUM]	
		GI:2222796	
3912	522	PROTEIN KINASE, PUTATIVE CONTAINS PFAM	Kinase, Protein
		PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE	į
		DOMAIN	
3919	523	DNA POLYMERASE III, EPSILON SUBUNIT GI:8163241	Polymerase
	*	FROM [CHLAMYDIA MURIDARUM][HYPOTHETICAL	Ì
		PROTEIN CONTAINS SIMILARITY TO]	
3923	524	ANTHRANILATE SYNTHASE LARGE SUBUNIT	Synthase
]		GI:1374671 FROM [BUCHNERA APHIDICOLA],	
2022		UNKNOWN PROTEIN CONTAINS SIMILARITY TO	
3932	525	GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
3952	526	CINNAMYL ALCOHOL DEHYDROGENASE, PUTATIVE	Dehydrogenases
	:	SIMILAR TO GB:X88797 FROM [EUCALYPTUS GUNNII]	
	İ	(PLANT MOL. BIOL. 36 (5), 755-765 (1998))	
3968	527	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE,	Transferases
		PUTATIVE SIMILAR TO PIR:T06460 FROM [PISUM	ļ
		SATIVUM]	
3983	528	NADP-SPECIFIC GLUTATAMATE DEHYDROGENASE,	Dehydrogenase
	1	PUTATIVE SIMILAR TO NADP-SPECIFIC	
		GLUTATAMATE DEHYDROGENASE (NADP-GDH)	
		SP:P28724 [GIARDIA LAMBLIA (GIARDIA	
		INTESTINALIS)]	
3986		REVERSE TRANSCRIPTASES[HYPOTHETICAL PROTEIN	l'ranscriptase
		SIMILAR TO]	
3987	530	IAA-ALA HYDROLASE (IAR3) IDENTICAL TO IAA-ALA	Hydrolase
	i	HYDROLASE (IAR3) [ARABIDOPSIS THALIANA]	
		GI:3421384	

3989	531	AUXIN CONJUGATE HYDROLASE (ILL5) IDENTICAL TO Hydrolase
		AUXIN CONJUGATE HYDROLASE [ARABIDOPSIS]
		THALIANA] (ILL5) GI:5725649;CONTAINS
		NONCONSENSUS AT ACCEPTOR SPLICE SITE AT
		EXON3
3990	532	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069: EUKARYOTIC PROTEIN
		KINASE DOMAIN, MULTIPLE PF00560: LEUCINE RICH
		REPEAT
3991	533	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069: EUKARYOTIC PROTEIN
		KINASE DOMAIN, MULTIPLE PF00560: LEUCINE RICH
		REPEAT
3992	534	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069: EUKARYOTIC PROTEIN
		KINASE DOMAIN, MULTIPLE PF00560: LEUCINE RICH
		REPEAT
3996	535	LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO LIGHT REPRESSIBLE
		RECEPTOR PROTEIN KINASE GI:1321686 FROM
		[ARABIDOPSIS THALIANA]
4001	536	LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO LIGHT REPRESSIBLE
		RECEPTOR PROTEIN KINASE GI:1321686 FROM
		[ARABIDOPSIS THALIANA]
4011	537	CAFFEIC ACID O-METHYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO CAFFEIC ACID O-METHYLTRANSFERASE
		GI:5031492 FROM [OCIMUM BASILICUM]
4039	538	PROTEIN KINASE, PUTATIVE SIMILAR TO GI:7573596 Kinase, Protein
		FROM [POPULUS NIGRA]
4041	539	PROTEIN KINASE, PUTATIVE SIMILAR TO GI:7573596 Kinase, Protein
		FROM [POPULUS NIGRA]
4044	540	SHORT CHAIN ALCOHOL DEHYDROGENASE, Dehydrogenase
		PUTATIVE SIMILAR TO GI:2739279 FROM [NICOTIANA
		TABACUM] (PLANT MOL. BIOL. 29 (5), 1027-1038 (1995))

4051	541	GLYCOSYL TRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		PFAM PROFILE: PF00534 GLYCOSYL TRANSFERASES	
		GROUP 1	
4053	542	GLYCOSYL TRANSFERASES-LIKE PROTEIN GI:9294599	Transferases
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO]	
4065	543	SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE,	Kinase, Protein
		PUTATIVE SIMILAR TO SOMATIC EMBRYOGENESIS	
		RECEPTOR-LIKE KINASE GI:2224910 FROM [DAUCUS	
	*	CAROTA]	
4081	544	LYSOPHOSPHOLIPASE GI:1552244 FROM [RATTUS	Lipase
		NORVEGICUS]; UNKNOWN PROTEIN CONTAINS	
		SIMILARITY TO	
4086	545		Hydrolase
4087	546	LIPASE SIMILAR TO MONOGLYCERIDE LIPASE	Lipase
		GB:NP_035974 FROM [MUS MUSCULUS][PUTATIVE]	
4090	547	OXYGENASE OXIDOREDUCTASE SIMILAR TO	Reductase
		ADVENTITIOUS ROOTING RELATED GB:CAA12386	
		FROM [MALUS DOMESTICA][PUTATIVE]	
4091	548		Oxidase
		[ARABIDOPSIS THALIANA][PUTATIVE],	
		OXIDOREDUCTASE SIMILAR TO	
4092	549		Reductase
		ADVENTITIOUS ROOTING RELATED GB:CAA12386	
4000		FROM [MALUS DOMESTICA][PUTATIVE]	
4093		GIBBERELLIN 20-OXIDASE GB:AAD42693 FROM	Oxidase
		[CITRULLUS LANATUS][PUTATIVE],	
4104	E E 1	OXIDOREDUCTASE SIMILAR TO	Dh h - 4
4104		PURPLE ACID PHOSPHATASE SIMILAR TO PURPLE	rnospnatase
		ACID PHOSPHATASE GB:CAA06921 FROM [IPOMOEA	
4110		BATATAS][PUTATIVE]	Camthoo
4110		DEOXYOCTULONOSIC ACID SYNTHETASE SIMILAR TO	Synthase
		3-DEOXY-MANNO-OCTULOSONATE  CYTIDYI VI TRANSEERASE CR. DOADS 1 FROM	
		CYTIDYLYLTRANSFERASE GB:P04951 FROM	
		[ESCHERICHIA COLI][PUTATIVE]	

4117	553	PROTEIN KINASE, PUTATIVE SIMILAR TO MANY Kinase, Protein
		PREDICTED PROTEIN KINASES
4118	554	PROTEIN KINASE, PUTATIVE SIMILAR TO MANY Kinase, Protein
		PREDICTED PROTEIN KINASES
4119	555	PROTEIN KINASE, PUTATIVE SIMILAR TO MANY Kinase, Protein
		PREDICTED PROTEIN KINASES
4143	556	PHOSPHOENOLPYRUVATE CARBOXYLASE 1, Carboxylase
		PUTATIVE SIMILAR TO PHOSPHOENOLPYRUVATE
[ [		CARBOXYLASE 1 GI:2266947 FROM [GOSSYPIUM
		HIRSUTUM]
4154	557	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE GB:AAC50043 GI:2465923
		FROM [ARABIDOPSIS THALIANA]
4155	558	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE GB:AAC50043 GI:2465923
		FROM [ARABIDOPSIS THALIANA]
4163	559	CHALCONE ISOMERASE, PUTATIVE SIMILAR TO Isomerase
		GI:4126399 FROM [CITRUS SINENSIS]
4178	560	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO Transferases
		GI:2853219 FROM [CARICA PAPAYA]
4183	561	LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN Kinase, Protein
		KINASE 1, PUTATIVE SIMILAR TO GI:3360289 FROM
		[ZEA MAYS] (PLANT MOL. BIOL. 37 (5), 749-761 (1998))
4184	562	LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN Kinase, Protein
		KINASE 1, PUTATIVE SIMILAR TO GI:3360289 FROM
l i		[ZEA MAYS] (PLANT MOL. BIOL. 37 (5), 749-761 (1998))
4193	563	PECTIN METHYLESTERASE (PMEU1), PUTATIVE Esterase
		SIMILAR TO PECTIN METHYLESTERASE (PMEU1)
		GI:1222551 FROM [LYCOPERSICON ESCULENTUM]
4194	564	PECTINESTERASE, PUTATIVE SIMILAR TO Esterase
		PECTINESTERASE GI:732912 FROM [PHASEOLUS
		VULGARIS]
		<del></del>

4202	565	LIPASE/HYDROLASE GDSL-like motif PROTEIN Lipase GI:9759145 FROM [ARABIDOPSIS THALIANA],
		UNKNOWN PROTEIN SIMILAR TO
4203	566	LIPASE, PUTATIVE SIMILAR TO LIPASE GI:1145627 Lipase
		FROM [ARABIDOPSIS THALIANA]
4208	567	LIPASE, PUTATIVE SIMILAR TO LIPASE GI:1145627 Lipase
		FROM [ARABIDOPSIS THALIANA]
4278	568	NADH DEHYDROGENASE SUBUNIT 3 GI:7542364 FROM Dehydrogenase
		[PAGURUS LONGICARPUS][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
4286	569	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN Kinase, Protein
		KINASE GI:9294282 FROM [ARABIDOPSIS THALIANA]
4298	570	GLYCOSYL TRANSFERASE GI:7021339 FROM Transferases
		[SINORHIZOBIUM MELILOTI][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
4299	571	NADH DEHYDROGENASE SUBUNIT 6 GI:2660574 FROM Dehydrogenase
		[FICEDULA HYPOLEUCA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
4300	572	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GI:3785984 FROM [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
4315	573	BETA-FRUCTOFURANOSIDASE, PUTATIVE SIMILAR TO Glycosylase
		BETA-FRUCTOFURANOSIDASE GI:9294027 FROM
		[ARABIDOPSIS THALIANA]
4323	574	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069 EUKARYOTIC PROTEIN KINASE
		DOMAIN
4332	575	LEUCOANTHOCYANIDIN DIOXYGENASE 2, PUTATIVE Oxygenases
		SIMILAR TO GI:5924383 FROM [DAUCUS CAROTA]
4351	576	TYROSINE PHOSPHATASE GB:AAF81798 GI:8926334 Phosphatase
		FROM [ORYZA SATIVA], SIMILAR TO PUTATIVE
4364	577	RECEPTOR KINASE, PUTATIVE SIMILAR TO CLVI Kinase, Protein
		RECEPTOR KINASE GB:AAB58929 GI:2160756 FROM
	i	[ARABIDOPSIS THALIANA]

4375	578	H+/CA2+ ANTIPORTER, PUTATIVE SIMILAR TO Transporter
		H+/CA2+ EXCHANGER 2 GB:BAA75232 GI:4512263 FROM
		[IPOMOEA NIL]
4376	579	H+/CA2+ ANTIPORTER, PUTATIVE SIMILAR TO Transporter
		H+/CA2+ EXCHANGER 2 GB:BAA75232 GI:4512263 FROM
		[IPOMOEA NIL]
4389	580	CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase
		PUTATIVE SIMILAR TO GB:AAD39534 FROM
		[GOSSYPIUM HIRSUTUM]
4396	581	SERINE ACETYLTRANSFERASE IDENTICAL TO Transferases
		GB:CAA84371 FROM [ARABIDOPSIS THALIANA] (EUR. J.
		BIOCHEM. 227 (1-2), 500-509 (1995))
4417	582	WALL-ASSOCIATED KINASE 2, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR-LIKE SERINE/THREONINE KINASE
		GB:AAC50043 GI:2465923 FROM [ARABIDOPSIS]
		THALIANA]
4418	583	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR Kinase, Protein
		TO RECEPTOR-LIKE SERINE/THREONINE KINASE
		GB:AAC50043 GI:2465923 FROM [ARABIDOPSIS
		THALIANA]
4419	584	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069: EUKARYOTIC PROTEIN
		KINASE DOMAIN, MULTIPLE PF00560: LEUCINE RICH REPEAT
4436	505	3'-5' EXONUCLEASE, PUTATIVE CONTAINS PFAM Nuclease
4430	303	PROFILE: PF01612: 3'-5' EXONUCLEASE
4464	586	TRIACYLGLYCEROL ACYLHYDROLASE GI:230348 Hydrolase
1704	200	FROM [RHIZOMUCOR MIEHEI][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
4469	587	CHITINASE, PUTATIVE SIMILAR TO CHITINASE Chitinase
		GI:17798 FROM [BRASSICA NAPUS]
4472	588	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase
		POLYGALACTURONASE PG1 GI:5669846 FROM
		[GLYCINE MAX]
4473	589	RECEPTOR-LIKE PROTEIN KINASE GI:7529754 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA], SIMILAR TO
		L

4480	590	PECTINACETYLESTERASE PRECURSOR, PUTATIVE E	sterase
		SIMILAR TO PECTINACETYLESTERASE PRECURSOR	
		GI:1431629 FROM [VIGNA RADIATA]	
4512	591	ATP PHOSPHORIBOSYL TRANSFERASE IDENTICAL TO T	ransferases
		ATP PHOSPHORIBOSYL TRANSFERASE GI:6683617	
		FROM [ARABIDOPSIS THALIANA]	
4518	592	PHOSPHOGLYCERATE KINASE, PUTATIVE SIMILAR TO K	inase
		PHOSPHOGLYCERATE KINASE GI:155584 FROM	
		[ZYMOMONAS MOBILIS]	
4520		CARBONIC ANHYDRASE, PUTATIVE SIMILAR TO A	nhydrase
	•	CARBONIC ANHYDRASE 1 GI:882241 FROM [FLAVERIA]	
		LINEARIS]	
4532	594	GLUTAMYL-TRNA REDUCTASE, PUTATIVE SIMILAR RO	eductase
		TO GLUTAMYL-TRNA REDUCTASE GI:1694925 FROM	
1500		[CUCUMIS SATIVUS]	
4533	595	HEME OXYGENASE PLASTID, PUTATIVE SIMILAR TO O	xygenases
		PLASTID HEME OXYGENASE GI:4877397 FROM	
4520	506	[ARABIDOPSIS THALIANA]	
4539	390	AMINO ACID PERMEASE I IDENTICAL TO AMINO ACID TE	ansporter
4541	507	PERMEASE I GI:22641 FROM [ARABIDOPSIS THALIANA]  XYLAN ENDOHYDROLASE, PUTATIVE SIMILAR TO H	udrologo
4341	391	(1,4)-BETA-XYLAN ENDOHYDROLASE GI:5306060 FROM	yuroiase
		[TRITICUM AESTIVUM]	
4545	508	PEPTIDYL-PROLYL ISOMERASE, PUTATIVE SIMILAR ISO	omerace
4545	370	TO PEPTIDYLPROLYL ISOMERASE GI:9294180 FROM	omerase
.		[ARABIDOPSIS THALIANA]	j
4553	599	XYLAN ENDOHYDROLASE ISOENZYME, PUTATIVE H	ydrolase
		SIMILAR TO XYLAN ENDOHYDROLASE ISOENZYME X-	
		I GI:1813594 FROM [HORDEUM VULGARE]	1
4571		GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Tr	ansferases
		TO GLUTATHIONE S-TRANSFERASE GB:AAF29773	
		GI:6856103 FROM [GOSSYPIUM HIRSUTUM]	ĺ
4573	601	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Tr	ansferases
		TO GLUTATHIONE S-TRANSFERASE GI:2190991 FROM	
		[AEGILOPS TAUSCHII]	
L		<u> </u>	

4578	602	OLICOPEDIDE TRANSPORTER BUTATIVE SIMILAR TO Transport
43/6	002	OLIGOPEPTIDE TRANSPORTER, PUTATIVE SIMILAR TO Transporter
		OLIGOPEPTIDE TRANSPORTER GI:510238 FROM
		[ARABIDOPSIS THALIANA]
4617	603	POTASSIUM TRANSPORTER, PUTATIVE SIMILAR TO Transporter
		POTASSIUM TRANSPORTER GI:2654088 FROM
•		[ARABIDOPSIS THALIANA]
4637	604	POLYGALACTURONASE ISOENZYME 1 BETA SUBUNIT, Glycosylase
		PUTATIVE SIMILAR TO POLYGALACTURONASE
		ISOENZYME 1 BETA SUBUNIT GI:1762585 FROM
]		[LYCOPERSICON ESCULENTUM]
4643	605	GALACTINOL SYNTHASE, PUTATIVE SIMILAR TO Synthase
		GALACTINOL SYNTHASE GI:5608497 FROM [AJUGA
		REPTANS]
4645	606	GALACTINOL SYNTHASE, PUTATIVE SIMILAR TO Synthase
		GALACTINOL SYNTHASE GI:5608497 FROM [AJUGA
		REPTANS]
4650	607	2-KETOCYCLOHEXANECARBOXYL-COA HYDROLASE Hydrolase
		GI:3243084 FROM [RHODOPSEUDOMONAS
		PALUSTRIS][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
4653	608	POLYGALACTURONASE, PUTATIVE SIMILAR TO Glycosylase
		POLYGALACTURONASE GI:9293940 FROM
		[ARABIDOPSIS THALIANA]
4655	609	S-RIBONUCLEASE BINDING PROTEIN SBP1 GI:6760451 Nuclease
1		FROM [PETUNIA HYBRIDA]
4657	610	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
		KINASE GI:498278 FROM [PETUNIA INTEGRIFOLIA]
4661	611	D-ALA,D-ALA LIGASE GI:6634594 FROM Ligase
		[STREPTOCOCCUS PNEUMONIAE][HYPOTHETICAL
		PROTEIN CONTAINS SIMILARITY TO]
4674	612	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein
	<del></del>	SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE
		GI:2224910 FROM [DAUCUS CAROTA]
4687	613	IRON-REGULATED TRANSPORTER SIMILAR TO IRON-Transporter
1007	013	REGULATED TRANSPORTER 1 GB:AAD30548 FROM
		[LYCOPERSICON ESCULENTUM][PUTATIVE]

4689	614	GIBBERELLIN 20-OXIDASE SIMILAR TO GIBBERELLIN Oxidase
		20-OXIDASE GB:CAA58295 FROM [ARABIDOPSIS
		THALIANA][PUTATIVE]
4690	615	AMINOMETHYLTRANSFERASE GB:CAA20175 FROM Transferases
		[STREPTOMYCES COELICOLOR A3(2)]; HYPOTHETICAL
		PROTEIN PREDICTED BY GENSCAN+, SIMILAR TO
4702	616	S-LINALOOL SYNTHASE, PUTATIVE SIMILAR TO S-Synthase
		LINALOOL SYNTHASE GI:1491939 FROM [CLARKIA
	•	BREWERI]
4715	617	SECRETORY CARRIER MEMBRANE PROTEIN Transporter
		IDENTICAL TO SECRETORY CARRIER MEMBRANE
		PROTEIN GI:7109228 FROM [ARABIDOPSIS THALIANA]
4717	618	LYSINE AND HISTIDINE SPECIFIC TRANSPORTER, Transporter
		PUTATIVE SIMILAR TO LYSINE AND HISTIDINE
		SPECIFIC TRANSPORTER GI:2576361 FROM
		[ARABIDOPSIS THALIANA]
4726	619	RECEPTOR PROTEIN KINASE (IRK1), PUTATIVE Kinase, Protein
		SIMILAR TO RECEPTOR PROTEIN KINASE (IRKI)
	<u>-</u>	GI:836953 FROM [IPOMOEA TRIFIDA]
4727	620	RECEPTOR PROTEIN KINASE (IRK1), PUTATIVE Kinase, Protein
		SIMILAR TO RECEPTOR PROTEIN KINASE (IRKI)
		GI:836953 FROM [IPOMOEA TRIFIDA]
4728	621	SERINE/THREONINE KINASE GI:4585880 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
4500		SIMILAR TO][PUTATIVE]
4729	622	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
4730	622	KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]  RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
4/30	023	KINASE 1 GI:2662048 FROM [BRASSICA RAPA]
4732	624	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
4/32	024	KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]
4733	625	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
41.23	023	KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]
4734	626	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
4/34	020	KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]
		MINIOU SI. 100072 I NOW [MANDIDOI SIS THADIAWA]

4736	627	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
		KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]	
4738	628	RECEPTOR PROTEIN KINASE (IRK1), PUTATIVE	Kinase, Protein
]		SIMILAR TO RECEPTOR PROTEIN KINASE (IRK1)	
		GI:836953 FROM [IPOMOEA TRIFIDA]	
4739	629	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
		KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]	
4740	630	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
		KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]	
4744	631	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
		KINASE GI:166692 FROM [ARABIDOPSIS THALIANA]	
4748	632	SERINE/THREONINE PROTEIN KINASE, PUTATIVE	Kinase, Protein
}		SIMILAR TO SERINE/THREONINE PROTEIN KINASE	
		GI:1066501 FROM [ARABIDOPSIS THALIANA]	
4750	633	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		RECEPTOR-LIKE KINASE GI:1783311 FROM [BRASSICA	
		OLERACEA]	
4752	634	INOSINE-GUANOSINE NUCLEOSIDE TRANSPORTER	Transporter
		GI:8272582 FROM [LEISHMANIA]	
]		DONOVANI][HYPOTHETICAL PROTEIN CONTAINS	
4756	625	SIMILARITY TO]  CINEOLE SYNTHASE, PUTATIVE SIMILAR TO 1,8-	Synthaga
4/36	033	CINEOLE SYNTHASE, FUTATIVE SIMILAR TO 1,6-	Syndiase
		OFFICINALIS]	
4760	636	DIHYDROFLAVONOL 4-REDUCTASE, PUTATIVE	Reductase
4700	350	SIMILAR TO DIHYDROFLAVONOL 4-REDUCTASE	l
		GI:1332411 FROM [ROSA HYBRIDA]	
4763	637	SERINE/THREONINE PROTEIN KINASE, PUTATIVE	Kinase, Protein
		SIMILAR TO SERINE/THREONINE PROTEIN KINASE	
	!	GI:3080385 FROM [ARABIDOPSIS THALIANA]	
4774	638	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE GI:9294282 FROM [ARABIDOPSIS THALIANA]	
4795	639	RECEPTOR-LIKE PROTEIN KINASE GI:3461841 FROM	Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
1		SIMILAR TO][PUTATIVE]	

4796	640	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR	Vinasa Destain
4790	040	TO RECEPTOR-LIKE PROTEIN KINASE, FOTATIVE SIMILAR	
		·	
		[ARABIDOPSIS THALIANA]	
4804	641	5'-ADENYLYLPHOSPHOSULFATE REDUCTASE,	
		PUTATIVE SIMILAR TO 5'-	
		ADENYLYLPHOSPHOSULFATE REDUCTASE GI:1336168	1
		FROM [ARABIDOPSIS THALIANA]	
4810	642	PEROXIDASE ISOZYME GI:217933 FROM [ARMORACIA	Oxidase
		RUSTICANA] UNKNOWN PROTEIN CONTAINS	
		SIMILARITY TO	
4823	643	ACC OXIDASE, PUTATIVE SIMILAR TO ACC OXIDASE	Oxidase
		GI:587086 FROM [BRASSICA OLERACEA]	
4843	644	GLUCOSE 1-DEHYDROGENASE (AB000617); SIMILAR	Dehydrogenases
		TO EST GB T88100 SIMILAR TO OXIDOREDUCTASE-	
		LIKE PROTEIN GB:CAB75763 GI:7019662 FROM	
		[ARABIDOPSIS THALIANA][SIMILAR TO]	
4846	645	3-KETOACYL-ACYL CARRIER PROTEIN SYNTHASE III	Synthase
		(KAS III) IDENTICAL TO 3-KETOACYL-ACYL CARRIER	
		PROTEIN SYNTHASE III (KAS III) GB:CAA72385	
		GI:1888359 [ARABIDOPSIS THALIANA]	
4863	646	AMINE OXIDASE, PUTATIVE SIMILAR TO AMINE	Oxidase
		OXIDASE GB:AAD49420 GI:5733089 [CANAVALIA	
		LINEATA]	
4871	647	NON-LTR REVERSE TRANSCRIPTASE, PUTATIVE	Transcriptase
		SIMILAR TO MANY PREDICTED NON-LTR REVERSE	•
		TRANSCRIPTASES	
4872	648	O-METHYLTRANSFERASE 1, PUTATIVE SIMILAR TO	Transferases
		GB:AAB96879 FROM [ARABIDOPSIS THALIANA]	
	į	(BIOCHIM. BIOPHYS. ACTA 1353 (3), 199-202 (1997))	
4877	649	PROTEIN KINASE, PUTATIVE CONTAINS PROTEIN	Kinase Protein
.0,,	0-17	KINASE DOMAINS	
4878	650		Synthase
70/0	050	SYNTHASE, PUTATIVE SIMILAR TO GB:U35779 FROM	Symmasc
		[TRITICUM AESTIVUM] (PLANT MOL. BIOL. 31 (5), 1009-	ĺ
		1020 (1996))	

4896	651	CAFFEIC O-METHYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO GI:602587 FROM [PRUNUS DULCIS] (PLANT
		PHYSIOL. 108, 1341-1341 (1995))
4921	652	GLUCOSE 1-DEHYDROGENASE GB:P40288 FROM Dehydrogenases
		[BACILLUS MEGATERIUM], REDUCTASE, PUTATIVE
		SIMILAR TO
4926	653	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069 EUKARYOTIC PROTEIN
		KINASE DOMAIN, PF00560 LEUCINE RICH REPEAT
4933	654	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069 EUKARYOTIC PROTEIN KINASE
	j	DOMAIN
4948	655	UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMATE Ligase
		2,6-DIAMINOPIMELATE LIGASE SIMILAR TO UDP-N-
		ACETYLMURAMOYLALANYL-D-GLUTAMATE2,6-
		DIAMINOPIMELATE LIGASE (MURE) GB:S40595
		[ESCHERICHIA COLI][PUTATIVE]
4956	656	AMINOPEPTIDASE SIMILAR TO AMINOPEPTIDASE N Protease
		(ALPHA-AMINOACYLPEPTIDE HYDROLASE) GB:P04825
		[ESCHERICHIA COLI]; CONTAINS PFAM PROFILE:
		PF00099 ZINC-BINDING METALLOPROTEASE
		DOMAIN[PUTATIVE]
4970	657	MONODEHYDROASCORBATE REDUCTASE SIMILAR TO Reductase
		MONODEHYDROASCORBATE REDUCTASE
		GB:AAD28178 [BRASSICA JUNCEA][PUTATIVE]
5024	658	PEPTIDE TRANSPORTER, PUTATIVE PREDICTED BY Transporter
		GENEMARK.HMM
5049	659	BETA-1,3-GLUCANASE, PUTATIVE SIMILAR TO BETA-Glycosylase
		1,3-GLUCANASE GI:8843743 FROM [ARABIDOPSIS
		THALIANA]
5059	660	RNA POLYMERASE SIGMA-SUBUNIT IDENTICAL TO Polymerase
		PLASTID RNA POLYMERASE SIGMA-SUBUNIT
	İ	GI:2398851 FROM [ARABIDOPSIS THALIANA]
5064	661	RHAMNOSYLTRANSFERASE, PUTATIVE SIMILAR TO Transferases
		RHAMNOSYLTRANSFERASE GI:454252 FROM [PETUNIA
		HYBRIDA]

5065	662	ANTHOCYANIDIN-3-GLUCOSIDE	Transferases
		RHAMNOSYLTRANSFERASE, PUTATIVE SIMILAR TO	
		ANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE GI:397567 FROM [PETUNIA	
		HYBRIDA]	
5070	663	GAMMA-TOCOPHEROL METHYLTRANSFERASE	Transferases
		ALMOST IDENTICAL TO GAMMA-TOCOPHEROL	
		METHYLTRANSFERASE GI:4106538 FROM	
		[ARABIDOPSIS THALIANA]	
5091	664	SERINE THREONINE KINASE GI:166813 FROM	Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		CONTAINS SIMILARITY TO]	
5096	665	ASPARTYL PROTEASE GI:6728988 FROM [ARABIDOPSIS	Protease
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	]
5097	666	SERINE THREONINE KINASE GI:166813 FROM	Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	!
		CONTAINS SIMILARITY TO]	
5103	667	XYLOGLUCAN ENDOTRANSGLYCOSYLASE, PUTATIVE	Glycosylase
		SIMILAR TO XYLOGLUCAN	
		ENDOTRANSGLYCOSYLASE I GI:3901012 FROM	
·		[FAGUS SYLVATICA]	
5114	668	ANTIGEN RECEPTOR, PUTATIVE SIMILAR TO ANTIGEN	Receptor
		RECEPTOR GI:3982955 FROM [GINGLYMOSTOMA	!
		CIRRATUM]	
5123	669	CARNITINE RACEMASE LIKE PROTEIN GI:7268149	Epimerase
		FROM [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO]	
5128	670	, and the second	Glycosylase
		POLYGALACTURONASE GI:2605891 FROM	
		[LYCOPERSICON ESCULENTUM]	
5129	671	INOSITOL POLYPHOSPHATE 5'-PHOSPHATASE	Phosphatase
		GI:3212848 FROM [ARABIDOPSIS THALIANA], SIMILAR	
		TO PUTATIVE	
5134	672	,	Protease
		PROTEASE DO (HTRA) SP:P09376 [ESCERICHIA COLI]	

5135	672	SERINE PROTEASE DO, PUTATIVE SIMILAR TO Protease
3133	0/3	
		PROTEASE DO (HTRA) SP:P09376 [ESCERICHIA COLI]
5150	674	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
		KINASE I [BRASSICA RAPA] GB:BAA23676
5151	675	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
		KINASE 1 [BRASSICA RAPA] GB:BAA23676
5166	676	GLUTAMATE DECARBOXYLASE (GAD), PUTATIVE Decarboxylase
1		SIMILAR TO GLUTAMATE DECARBOXYLASE (GAD)
		GI:294111 FROM [PETUNIA HYBRIDA]
5171	677	TERPENE SYNTHASE, PUTATIVE CONTAINS PFAM Synthase
	İ	PROFILE: PF01397: TERPENE SYNTHASE FAMILY
5182	678	OXIDOREDUCTASE, PUTATIVE CONTAINS PFAM Reductase
	]	PROFILE: PF01408: OXIDOREDUCTASE, GFO/IDH/MOCA
		FAMILY
5184	679	RECEPTOR PROTEIN KINASE (TMK1), PUTATIVE Kinase, Protein
		SIMILAR TO PUTATIVE RECEPTOR PROTEIN KINASE
		(TMK1) [ARABIDOPSIS THALIANA (MOUSE-EAR
		CRESS)] SP:P43298
5194	680	BETA-1,3-GLUCANASE PRECURSOR, PUTATIVE Glycosylase
	į	SIMILAR TO GI:4097948 FROM [ORYZA SATIVA] (GENE
		223 (1-2), 311-320 (1998))
5212	681	FRUCTOKINASE, PUTATIVE SIMILAR TO Kinase
		FRUCTOKINASE [LYCOPERSICON ESCULENTUM]
		GI:2102691
5215	682	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE
		DOMAIN
5221	683	FORMYL TRANSFERASE, PUTATIVE CONTAINS PFAM Transferases
		PROFILE: PF00551: FORMYL TRANSFERASE
5226		SUCROSE-PROTON SYMPORTER, 5' PARTIAL SIMILAR Transporter
		TO C-TERM OF SUCROSE-PROTON SYMPORTER
		[ARABIDOPSIS THALIANA] GI:407094
5248		CINNAMYL ALCOHOL DEHYDROGENASE, PUTATIVE Dehydrogenases
3246		SIMILAR TO CINNAMYL ALCOHOL DEHYDROGENASE
		[EUCALYPTUS GUNNII] GI:1143445

	50.6	Propress of the control of the contr
5251	686	RECEPTOR PROTEIN KINASE, PUTATIVE CONTAINS Kinase, Protein
		PFAM PROFILES: PF00069: EUKARYOTIC PROTEIN
		KINASE DOMAIN, MULTIPLE PF00560: LEUCINE RICH
		REPEAT
5256	687	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
		PROFILE: PF00069: EUKARYOTIC PROTEIN KINASE
		DOMAIN
5259	688	RECEPTOR SERINE/THREONINE KINASE PR5K, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR
[		SERINE/THREONINE KINASE PR5K GI:1235680 FROM
		[ARABIDOPSIS THALIANA]
5260	689	RECEPTOR SERINE/THREONINE KINASE PR5K, Kinase, Protein
) j		PUTATIVE SIMILAR TO RECEPTOR
		SERINE/THREONINE KINASE PR5K GI:1235680 FROM
		[ARABIDOPSIS THALIANA]
5261	690	RECEPTOR SERINE/THREONINE KINASE PR5K, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR
		SERINE/THREONINE KINASE PR5K GI:1235680 FROM
		[ARABIDOPSIS THALIANA]
5266	691	KINASE, PUTATIVE SIMILAR TO LEAF RUST Kinase, Protein
		RESISTANCE KINASE LR10 GI:1680685 FROM
		[TRITICUM AESTIVUM]
5267	692	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR-LIKE KINASE GI:5523856 FROM [HORDEUM
		VULGARE]
5275	693	RIBULOSE BISPHOSPHATE CARBOXYLASE SMALL Carboxylase
		UNIT, PUTATIVE SIMILAR TO RIBULOSE-
		BISPHOSPHATE CARBOXYLASE SMALL UNIT GI:406726
		FROM [BRASSICA NAPUS]
5292	694	GLYOXAL OXIDASE (GLX1), PUTATIVE SIMILAR TO Oxidase
- 1		GLYOXAL OXIDASE (GLX1) GI:1050301 FROM
		[PHANEROCHAETE CHRYSOSPORIUM]
5309	695	PROTEIN KINASE PFAM HMM HIT: EUKARYOTIC Kinase, Protein
	0,5	PROTEIN KINASE DOMAIN; IDENTICAL TO
		GB:AAC18787 [ARABIDOPSIS THALIANA][PUTATIVE]
<u></u> 1		OD. ANGLOVO [ALADIDOI SIG THADIAWA][FOTATIVE]

5313	696	RECEPTOR-LIKE PROTEIN KINASE GI:4262228 FROM	Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
		SIMILAR TO][PUTATIVE]	
5314	697	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR	Kinase, Protein
		KINASE 1 GI:9294449 FROM [ARABIDOPSIS THALIANA]	·
5325	698	AMINO ACID PERMEASE SIMILAR TO LYSINE AND	Transporter
		HISTIDINE SPECIFIC TRANSPORTER GB:AAC49885	
		[ARABIDOPSIS THALIANA][PUTATIVE]	
5330	699	AMINO ACID PERMEASE, PUTATIVE SIMILAR TO	Transporter
		AMINO ACID PERMEASE 1 GB: AAB48944 GI:976402	
		FROM [NICOTIANA SYLVESTRIS]	
5341	700	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
		LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE	
		GB:CAA66376 GI:1321686 FROM [ARABIDOPSIS	
		THALIANA]	
5344	701	ENDOCHITINASE, PUTATIVE SIMILAR TO PECTATE	Chitinase
		LYASE 1 GB:AAF19195 GI:6606532 FROM [MUSA	
		ACUMINATA]	
5346	702	ABC TRANSPORTER SIMILAR TO TERMINAL EARI	Transporter
		GB:AAC39463 GI:3153237 FROM [ZEA MAYS] (NATURE	
		393 (6681), 166-168 (1998))[PUTATIVE]	
5358	703		Transferases
		GB:AAC53064 [MUS MUSCULUS], UNKNOWN PROTEIN	İ
-2.60	~~.	SIMILAR TO	
5368	704		Transferases
		COENZYME A 3-O-METHYLTRANSFERASE SIMILAR TO	
		S-ADENOSYL-L-METHIONINE:TRANS-CAFFEOYL- COENZYME A 3-O-METHYLTRANSFERASE	
		GB:AAA62426 [ARABIDOPSIS THALIANA]	
		(FUNCTION=DISEASE RESISTANCE)[PUTATIVE]	
5369	705	1	Transferases
3309	703	COENZYME A 3-O-METHYLTRANSFERASE SIMILAR TO	1101131010305
		S-ADENOSYL-L-METHIONINE:TRANS-CAFFEOYL-	
}		COENZYME A 3-O-METHYLTRANSFERASE	
		GB:AAA62426 [ARABIDOPSIS THALIANA]	
		(FUNCTION=DISEASE RESISTANCE)[PUTATIVE]	
		[. J. J. J. J. J. J. J. J. J. J. J. J. J.	

5370	706	PHOSPHATIDYLINOSITOL SYNTHASE (PIS1) Synthase
		IDENTICAL TO PHOSPHATIDYLINOSITOL SYNTHASE
		(PIS1) GB:AJ000539
5371	707	HYDROXYPYRUVATE REDUCTASE (HPR) IDENTICAL Reductase
		TO HYDROXYPYRUVATE REDUCTASE (HPR)
		GB:D85339 [ARABIDOPSIS THALIANA] (PLANT CELL
		PHYSIOL 1997 APR;38(4):449-55)
5374	708	S-ADENOSYL-L-METHIONINE:SALICYLIC ACID Transferases
}		CARBOXYL METHYLTRANSFERASE SIMILAR TO S-
]		ADENOSYL-L-METHIONINE:SALICYLIC ACID
		CARBOXYL METHYLTRANSFERASE GB:AAF00108
		[CLARKIA BREWERI] (ENZYME INVOLVED IN FLORAL
		SCENT PRODUCTION AND PLANT
		DEFENSE)[PUTATIVE]
5399	709	BIFUNCTIONAL NUCLEASE SIMILAR TO Nuclease
		BIFUNCTIONAL NUCLEASE GB:AAD00694 FROM
		[ZINNIA ELEGANS][PUTATIVE]
5410	710	RECEPTOR KINASE SIMILAR TO RECEPTOR KINASE Kinase, Protein
		GB:AAA33715 FROM [PETUNIA]
5411	711	INTEGRIFOLIA][PUTATIVE]
5411	/11	PROTEIN PHOSPHATASE SIMILAR TO PROTEIN Phosphatase PHOSPHATASE-2C GB:AAC36697 FROM
		[MESEMBRYANTHEMUM CRYSTALLINUM][PUTATIVE]
5412	712	ASPARAGINYL-TRNA SYNTHETASE SIMILAR TO Synthase
3412	/12	SYNC1 PROTEIN GB:AAD46681[PUTATIVE]
5416	713	TRNA ISOPENTENYL TRANSFERASE SIMILAR TO TRNA Transferases
3,10	713	ISOPENTENYL TRANSFERASE GB:AAF00582
		[ARABIDOPSIS THALIANA]; CONTAINS PFAM PROFILE:
		PF01715 IPP TRANSFERASE (TRNA DELTA(2)-
		ISOPENTENYLPYROPHOSPHATE
		TRANSFERASE)[PUTATIVE]
5424	714	CINNAMOYL COA REDUCTASE GB:CAA12276 Reductase
		[POPULUS BALSAMIFERA SUBSP. TRICHOCARPA],
		ALDEHYDE REDUCTASE GB:AAD53967 [VIGNA
	!	RADIATA], DIHYDROFLAVONOL 4-REDUCTASE
		GB:BAA12723 [ROSA HYBRIDA][PUTATIVE]
L		l

5439	715	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase Protein
		KINASE 1 GB:BAA94509 GI:7573596 FROM [POPULUS	1
		NIGRA]	!
5441	716	·	ATD
5441	/16	ATPASE, PUTATIVE SIMILAR TO CHROMAFFIN	ļ
]	l	GRANULE ATPASE II GB:AAD03352 GI:4115341 FROM	
	<u> </u>	[BOS TAURUS]	
5442	717	CYTIDINE AND DEOXYCYTIDYLATE DEAMINASE	Deaminase
j	j	ZINC-BINDING REGION [UNKNOWN PROTEIN	
		CONTAINS PFAM PROFILE: PF00383]	
5446	718	PHOSPHOENOLPYRUVATE CARBOXYLASE SIMILAR	Carboxylase
ļ		TO PHOSPHOENOLPYRUVATE CARBOXYLASE	
	•	GB:CAA11415 [BRASSICA JUNCEA]; CONTAINS PFAM	
		PROFILE: PF00311 PHOSPHOENOLPYRUVATE	
		CARBOXYLASE[PUTATIVE]	
5453	719	PROTEIN KINASE PFAM HMM HIT: EUKARYOTIC	Kinase, Protein
	}	PROTEIN KINASE DOMAIN[PUTATIVE]	
5455	720	PEROXIDASE ATP23A IDENTICAL TO PEROXIDASE	Oxidase
		ATP23A GB:CAA70035 [ARABIDOPSIS THALIANA]	
5458	721		Synthase
		CARBOXYLIC ACID SYNTHASE / 2-OXOGLUTARATE	
		DECARBOXYLASE, MENAQUINONE BIOSYNTHESIS	
		PROTEIN SIMILAR TO MENAQUINONE BIOSYNTHESIS	
ļ		PROTEIN GB:P23970 [BACILLUS SUBTILIS]; PFAM HMM	
	ļ	HIT: [PUTATIVE]	
5492	722		V:
5482	122	DIHYDROPTERIN PYROPHOSPHOKINASE, PUTATIVE	Kinase
		SIMILAR TO DIHYDROPTERIN PYROPHOSPHOKINASE	
		GI:1934971 FROM [PISUM SATIVUM]	
5483	723	FRUCTOKINASE SIMILAR TO FRUCTOKINASE	Kinase
	}	GB:AAB57733 [LYCOPERSICON]	
		ESCULENTUM][PUTATIVE]	
5487	724	POLYNEURIDINE ALDEHYDE ESTERASE, PUTATIVE	Esterase
		SIMILAR TO POLYNEURIDINE ALDEHYDE ESTERASE	
		GI:6651393 FROM [RAUVOLFIA SERPENTINA]	
5490	725	RECEPTOR-LIKE PROTEIN KINASE SIMILAR TO	Kinase, Protein
		PROTEIN KINASE GB:AAD11518 [ARABIDOPSIS	j
		THALIANA][PUTATIVE]	
1		·	ł

5499	726	CHORISMATE MUTASE, PUTATIVE CONTAINS PFAM Mutase
		PROFILE: PF01817: CHORISMATE MUTASE
5525	727	STEROL DESATURASE CONTAINS PFAM PROFILE: Desaturases
		PF01598 STEROL DESATURASE[PUTATIVE]
5533	728	HEME OXYGENASE SIMILAR TO HEME OXYGENASE Oxygenases
		GB:AAD22107 [ARABIDOPSIS THALIANA][PUTATIVE]
5534	729	PROTEIN KINASE CONTAINS PFAM PROFILE: PF00069 Kinase, Protein
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
5544	730	ALPHA-AMYLASE SIMILAR TO ALPHA-AMYLASE Glycosylase
		GB:AAA91884 [SOLANUM TUBEROSUM][PUTATIVE]
5546	731	NITRATE TRANSPORTER (NTL1) IDENTICAL TO Transporter
		NITRATE TRANSPORTER (NTL1) GB:AAC28086
		[ARABIDOPSIS THALIANA]
5547	732	PEPTIDE TRANSPORTER SIMILAR TO PEPTIDE Transporter
		TRANSPORTER GB:AAC32034 [HORDEUM
		VULGARE][PUTATIVE]
5548	733	PEPTIDE TRANSPORTER SIMILAR TO PEPTIDE Transporter
		TRANSPORTER GB:AAC32034 [HORDEUM
7.5.50		VULGARE][PUTATIVE]
5552		PROTEIN KINASE PFAM HMM HIT: EUKARYOTIC Kinase, Protein
5550		PROTEIN KINASE DOMAIN[PUTATIVE]
5553	735	GLUTATHIONE TRANSFERASE SIMILAR TO Transferases
		GLUTATHIONE TRANSFERASE GB:CAA09188 [ALOPECURUS MYOSUROIDES][PUTATIVE]
5554		GLUTATHIONE TRANSFERASE SIMILAR TO Transferases
3334	750	GLUTATHIONE TRANSFERASE GB:CAA09188
		[ALOPECURUS MYOSUROIDES][PUTATIVE]
5555		PECTIN METHYLESTERASE SIMILAR TO PECTIN Esterase
		METHYLESTERASE GB:CAA96435 [NICOTIANA]
	j	PLUMBAGINIFOLIA][PUTATIVE]
5560		RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR Kinase, Protein
		TO RECEPTOR-LIKE PROTEIN KINASE GI:8777368 FROM
	ł	[ARABIDOPSIS THALIANA]
5568		VETISPIRADIENE SYNTHASE GI:9294376 FROM Synthase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
	· · · · · · · · · · · · · · · · · · ·	SIMILAR TO]

5569	740	GLYCOSYL TRANSFERASE GI:7268597 FROM Transferases
		[ARABIDOPSIS THALIANA], UNKNOWN PROTEIN
		SIMILAR TO PUTATIVE
5571	741	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR-LIKE KINASE GI:1405837 FROM
		[ARABIDOPSIS THALIANA]
5573	742	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR-LIKE KINASE GI:1405837 FROM
		[ARABIDOPSIS THALIANA]
5585	743	RECEPTOR SERINE/THREONINE KINASE PR5, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR
		SERINE/THREONINE KINASE PR5 GI:1235680 FROM
		[ARABIDOPSIS THALIANA]
5590	744	POTASSIUM TRANSPORTER, PUTATIVE SIMILAR TO Transporter
		POTASSIUM TRANSPORTER GI:2654088 FROM
		[ARABIDOPSIS THALIANA]
5601	745	CARBONIC ANHYDRASE, PUTATIVE SIMILAR TO Anhydrase
		CARBONIC ANHYDRASE GI:882241 FROM [FLAVERIA
		LINEARIS]
5605	746	PROTEIN KINASE SIMILAR TO PROTEIN KINASE Kinase, Protein
		(APK1A) GB:Q06548 [ARABIDOPSIS THALIANA],
		IDENTICAL TO RESIDUES 1-319 OF SERINE/THREONINE
		KINASE-LIKE PROTEIN GB:AAC18796 [ARABIDOPSIS
		THALIANA]; PFAM HMM HIT: EUKARYOTIC PROTEIN
5506	= 45	KINASE DOMAIN[PUTATIVE]
5606	i I	PROTEIN KINASE SIMILAR TO C-TERMINAL REGION Kinase, Protein
		HAS SIMILARITY TO C-TERMINAL REGION OF
		PROTEIN KINASE (APK1A) GB:Q06548 [ARABIDOPSIS
		THALIANA]; PFAM HMM HIT: EUKARYOTIC PROTEIN
5610	710	KINASE DOMAIN[PUTATIVE]  POLYGALACTURONASE SIMILAR TO Glycosylase
2010		POLYGALACTURONASE SIMILAR TO Glycosylase POLYGALACTURONASE PRECURSOR (PECTINASE)
		` '
5612	740	GB:P35336 [ACTINIDIA CHINENSIS][PUTATIVE] PROTEIN KINASE SIMILAR TO C-TERMINAL REGION Kinase, Protein
3012	147	OF S-RECEPTOR KINASES GB:BAA21132, GB:BAA06285
		[BRASSICA RAPA]; PFAM HMM HIT: EUKARYOTIC
		PROTEIN KINASE DOMAIN[PUTATIVE]
		. NO LEM KINASE DOMAIN[FUTATIVE]

5613	750	PROTEIN KINASE SIMILAR TO C-TERMINAL REGION	Kinase, Protein
		OF S-RECEPTOR KINASE PRECURSOR GB:JQ1677	
		[BRASSICA NAPUS]; PFAM HMM HIT: EUKARYOTIC	
		PROTEIN KINASE DOMAIN[PUTATIVE]	
5634	751	PROTEIN KINASE CONTAINS PFAM PROFILE: PF00069	Kinase, Protein
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]	
5642	752	PHOSPHOGLUCOMUTASE, PUTATIVE SIMILAR TO	Mutase
		PHOSPHOGLUCOMUTASE GI:534981 FROM [SPINACIA	
		OLERACEA]	
5653	753	AUXIN TRANSPORT PROTEIN REH1, PUTATIVE	Transporter
		SIMILAR TO AUXIN TRANSPORT PROTEIN REHI	
		GI:3377509 FROM [ORYZA SATIVA]	
5663	754	BILIRUBIN OXIDASE SIMILAR TO; SPORE COAT	Oxidase
		PROTEIN-LIKE PROTEIN SIMILAR TO SPORE COAT	
		PROTEIN A GB:P07788 FROM [BACILLUS SUBTILIS]	
		AND GB:Q12737 [MYROTHECIUM VERRUCARIA]	
5666	755	GLUCOSAMINYL (N-ACETYL) TRANSFERASE	Transferases
		GB:4758422 FROM [HOMO SAPIENS], UNKNOWN	ĺ
		PROTEIN SIMILAR TO	
5675	756	FATTY ACID ELONGASE 3-KETOACYL-COA SYNTHASE	Synthase
		i, ketoacyl-coa synthase similar to,	
		GB:AAC99312[PUTATIVE]	
5676	757	DEHYDROGENASE SIMILAR TO D-THREONINE	Dehydrogenases
		DEHYDROGENASE GB:BAA34184[PUTATIVE]	
5677	758	DEHYDROGENASE SIMILAR TO D-THREONINE	Dehydrogenases
		DEHYDROGENASE GB:BAA34184[PUTATIVE]	
5681	759	·	Transferases
		GLUCOSYLTRANSFERASE SIMILAR TO UDP-	
		GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE	
		PRECURSOR GB:Q09332 [DROSOPHILA	
5694	260	MELANOGASTER][PUTATIVE]	Y :
5684	/60	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE CONTAINS PFAM PROFILE: PF00657	Lipase
5712	761	LIPASE/ACYLHYDROLASE  AMINO ACID PERMEASE SIMILAR TO AMINO ACID	Transporter
3/12			Tansporter
]		PERMEASE GB:AAB48944 FROM [NICOTIANA SYLVESTRIS][PUTATIVE]	
		21 LA C21 KIO][LÛ IA IIAC]	

5714	762	LIPASE/HYDROLASE GDSL-motif SIMILAR TO Lipase
		PUTATIVE LIPASE/HYDROLASE GDSL-
		motifGB:AAC23769 AND GB:AAD12024[PUTATIVE]
5716	763	PEROXIDASE ATP4A IDENTICAL TO GB:CAA67309 Oxidase
	•	GI:1429213 FROM [ARABIDOPSIS THALIANA]
5723	764	HYPOXANTHINE RIBOSYL TRANSFERASE, PUTATIVE Transferases
		SIMILAR TO HYPOXANTHINE RIBOSYL TRANSFERASE
		GB:AAC46403 GI:2689037 FROM [VIBRIO
		PARAHAEMOLYTICUS]
5731	765	PROTEIN KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
		PROTEIN KINASE GB:BAA11869 GI:1389566 FROM
		[ARABIDOPSIS THALIANA]
5736	766	SUCROSE TRANSPORT PROTEIN SUC1 IDENTICAL TO Transporter
7707		GB:S38197 FROM [ARABIDOPSIS THALIANA]
5737	767	SUCROSE TRANSPORT PROTEIN SIMILAR TO Transporter
		SUCROSE TRANSPORT PROTEIN SUCI GB:S38197
5740	760	FROM [ARABIDOPSIS THALIANA][PUTATIVE] HISTIDINOL-PHOSPHATE AMINOTRANSFERASE Transferases
3/40	708	SIMILAR TO HISTIDINOL-PHOSPHATE
		AMINOTRANSFERASE GB:CAA70403 FROM
		[NICOTIANA TABACUM][PUTATIVE]
5747	769	FUCOSYLTRANSFERASE 9 GB:NP_034373 FROM [MUS Transferases
		MUSCULUS], HYPOTHETICAL PROTEIN PREDICTED
		BY GENEFINDER, CONTAINS SIMILARITY TO
5761	770	OLIGOPEPTIDE TRANSPORTER, PUTATIVE SIMILAR TO Transporter
		LEOPT1 [LYCOPERSICON ESCULENTUM] GI:4102839
5766	771	LEUCINE-RICH RECEPTOR-LIKE PROTEIN KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO GI:3641252 FROM [MALUS X
		DOMESTICA] (PLANT MOL. BIOL. 40 (6), 945-957 (1999))
5767	772	PHOSPHOGLYCERATE DEHYDROGENASE, PUTATIVE Dehydrogenase
		CONTAINED WITHIN GI:6626247 FROM
		[ARCHAEOGLOBUS FULGIDUS]
5777	773	LEUCINE-RICH RECEPTOR-LIKE PROTEIN KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO GI:3641252 FROM [MALUS X
		DOMESTICA] (PLANT MOL. BIOL. 40 (6), 945-957 (1999))

5793	774	RECEPTOR-LIKE PROTEIN KINASE CONTAINS PFAM Kinas	se, Protein
		PROFILES: PF00560 LEUCINE RICH REPEAT (5 COPIES),	
		PF00069 EUKARYOTIC PROTEIN KINASE	
		DOMAIN[PUTATIVE]	
5801	775	PROTEIN KINASE SIMILAR TO PROTEIN KINASE APK1A Kinas	e, Protein
		GB:Q06548 [ARABIDOPSIS THALIANA][PUTATIVE]	
5815	776	CINNAMYL-ALCOHOL DEHYDROGENASE SIMILAR TO Dehy	drogenases
ĺ		CINNAMYL-ALCOHOL DEHYDROGENASE	
		GB:AAC35846 [MEDICAGO SATIVA][PUTATIVE]	
5817	777	P-TYPE TRANSPORTING ATPASE SIMILAR TO ATPASE Trans	porter
		II GB:AAD34706 [HOMO SAPIENS][PUTATIVE]	
5827	778	THREONINE SYNTHASE, PUTATIVE SIMILAR TO Synth	ase
		THREONINE SYNTHASE GI:4850369 FROM	
70.10		[ARABIDOPSIS THALIANA]	
5843	779	MANDELONITRILE LYASE GI:288115 FROM [PRUNUS Lyase	
		SEROTINA], UNKNOWN PROTEIN CONTAINS	
5851	790	SIMILARITY TO  MANDELONITRILE LYASE, PUTATIVE SIMILAR TO Lyase	
3631	760	MANDELONITRILE LYASE GI:288115 FROM [PRUNUS	
		SEROTINA]	
5862	781	GLYCOSYL TRANSFERASE CONTAINS PFAM PROFILE: Transi	ferases
		PF00534 GLYCOSYL TRANSFERASES GROUP	
		I[PUTATIVE]	
5880	782	STEROID 22-ALPHA-HYDROXYLASE, PUTATIVE Hydro	xylase
		SIMILAR TO STEROID 22-ALPHA-HYDROXYLASE	
		GI:2935342 FROM [ARABIDOPSIS THALIANA]	
5881	783	GLUCOSIDASE II BETA-SUBUNIT GI:5452942 FROM Glyco	sylase
		[MUS MUSCULUS][HYPOTHETICAL PROTEIN	-
		CONTAINS SIMILARITY TO]	
5883	784	SUCROSE SYNTHASE, PUTATIVE SIMILAR TO Syntha	ase
		SUCROSE SYNTHASE GI:4468151 FROM	
		[CRATEROSTIGMA PLANTAGINEUM]	
5893	785	LYSOPHOSPHOLIPASE HOMOLOG, PUTATIVE SIMILAR Lipase	;
		TO LYSOPHOSPHOLIPASE HOMOLOG GI:2801536 FROM	
		[ORYZA SATIVA]	

5898	786	DIPHOSPHOINOSITOL POLYPHOSPHATE Hydrolase
3070	/00	PHOSPHOHYDROLASE GI:3978224 FROM [HOMO]
		SAPIENS], UNKNOWN PROTEIN CONTAINS
		SIMILARITY TO
5903	707	
3903	/6/	AUXIN TRANSPORTER SPLICE VARIANT B, PUTATIVE Transporter
		SIMILAR TO GI:7109715 FROM [ARABIDOPSIS
5905	700	THALIANA]
3903	/00	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
5010	700	LIPASE/ACYLHYDROLASE
5918	/89	UDP-N-ACETYLGLUCOSAMINEN-ACETYLMURAMYL- Transferases
		(PENTAPEPTIDE)-PYROPHOSPHORYL-UNDECAPRENOL
		N-ACETYLGLUCOSAMINE TRANSFERASE SIMILAR TO
		PUTATIVE UDP-N-ACETYLGLUCOSAMINEN-
		ACETYLMURAMYL-(PENTAPEPTIDE)-
		PYROPHOSPHORYL-UNDECAPRENOL N-
		ACETYLGLUCOSAMINE TRANSFERASE GB:007670
5919	700	[ENTEROCOCCUS HIRAE][PUTATIVE]  ALPHA/BETA HYDROLASE FOLD  Hydrolase
5932		ALPHA/BETA HYDROLASE FOLD Hydrolase GLUCOSYLTRANSFERASE SIMILAR TO Transferases
3932	791	GLUCOSYLTRANSFERASE GB:AAB36653 FROM
5935	702	[NICOTIANA TABACUM][PUTATIVE]  LIPASE SIMILAR TO LIPASE GB:CAA74737 FROM Lipase
3933	192	[DROSOPHILA MELANOGASTER][PUTATIVE]
5941	702	URIDINE KINASE GB:AAF09747 FROM [DEINOCOCCUS Kinase
3941	193	
		RADIODURANS], UNKNOWN PROTEIN CONTAINS A REGION SIMILAR TO
5040	704	
5942		PROTEASE IV SIMILAR TO PROTEASE IV GB:AAA57008 Protease
5042		FROM [ESCHERICHIA COLI][PUTATIVE]
5943	195	STRICTOSIDINE SYNTHASE SIMILAR TO Synthase
		STRICTOSIDINE SYNTHASE 3 PRECURSOR
5044	70.5	GB:P92976[PUTATIVE]
5944	796	STRICTOSIDINE SYNTHASE SIMILAR TO Synthase
		STRICTOSIDINE SYNTHASE 1/2 PRECURSOR
		GB:P94111[PUTATIVE]

5945	. 797	STRICTOSIDINE SYNTHASE EXTREMELY SIMILAR TO Synthase
		STRICTOSIDINE SYNTHASE 1/2 PRECURSOR
]		GB:P94111[PUTATIVE]
5947	798	2-ISOPROPYLMALATE SYNTHASE SIMILAR TO 2-Synthase
		ISOPROPYLMALATE SYNTHASE GB: AAB61598 FROM
		[LYCOPERSICON PENNELLII][PUTATIVE]
5950	799	ADENOSINE KINASE (EC 2.7.1.20) GB:S52758 FROM Kinase
		[LEISHMANIA DONOVANI], HYPOTHETICAL PROTEIN
		PREDICTED BY GENSCAN+, SIMILAR TO
5952	800	FLAVONOL SULFOTRANSFERASE SIMILAR TO Transferases
		FLAVONOL 4'-SULFOTRANSFERASE GB:P52837 FROM
		[FLAVERIA CHLORAEFOLIA][PUTATIVE]
5964	801	GLYCEROPHOSPHODIESTER PHOSPHODIESTERASE Esterase
		SIMILAR TO GLYCEROPHOSPHORYL DIESTER
		PHOSPHODIESTERASE GB:AAF12F49 FROM
		[DEINOCOCCUS RADIODURANS][PUTATIVE]
5967	802	MITOCHONDRIAL CARRIER PROTEIN SIMILAR TO Transporter
		GB:CAA21437 FROM [SCHIZOSACCHAROMYCES]
		POMBE][PUTATIVE]
5979	803	RECEPTOR PROTEIN KINASE SIMILAR TO Kinase, Protein
		BRASSINOSTEROID INSENSITIVE 1 GB:AAC49810
		(PUTATIVE RECEPTOR PROTEIN KINASE); CONTAINS
		PFAM PROFILES: PF00560 LEUCINE RICH REPEAT (17
		REPEATS), PF00069 EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
5981	804	ALPHA GALACTOSYLTRANSFERASE SIMILAR TO Transferases
3901	004	ALPHA GALACTOSYLTRANSFERASE GB:CAB52246
		[TRIGONELLA FOENUM-GRAECUM] (PLANT CELL
		WALL MATRIX POLYSACCHARIDE
		BIOSYNTHESIS)[PUTATIVE]
5982	805	EXONUCLEASE CONTAINS PFAM PROFILE: PF00929 Nuclease
		EXONUCLEASE[PUTATIVE]
5985	806	XYLOGLUCAN FUCOSYLTRANSFERASE SIMILAR TO Transferases
		XYLOGLUCAN FUCOSYLTRANSFERASE GB:AAD41092
		[ARABIDOPSIS THALIANA] (ENZYME INVOLVED IN
		PLANT CELL WALL BIOSYNTHESIS: SCIENCE 284, 1976-
		1979 (1999))[PUTATIVE]
L		

5989	807	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE	Lipase
		CONTAINS PFAM PROFILE: PF00657	
		LIPASE/ACYLHYDROLASE	
5990	808	GERANYLGERANYL REDUCTASE IDENTICAL TO	Reductase
		GERANYLGERANYL REDUCTASE GB:Y14044	
		[ARABIDOPSIS THALIANA] (INVOLVEMENT:	
		CHLOROPHYLL, THE TOCOPHEROL AND THE	
		PHYLLOQUINONE PATHWAYS EUR J BIOCHEM 1998	
į	}	JAN 15;251(1-2):413-7)	
6001	809	GLUTATHIONE S-TRANSFERASE SIMILAR TO	Transferases
		PUTATIVE GLUTATHIONE S-TRANSFERASE	
		GB:CAA10060 [ARABIDOPSIS THALIANA]; CONTAINS	
		PFAM PROFILE: PF00043 GLUTATHIONE S-	
		TRANSFERASES[PUTATIVE]	
6005	810	ALPHA/BETA HYDROLASE FOLD	Hydrolase
6012	811	ISOCHORISMATE SYNTHASE (ICSI) IDENTICAL TO	Synthase
		ISOCHORISMATE SYNTHASE (ICSI) GB:AF078080	
		(CATALYZES CHORISMIC ACID TO ISOCHORISMIC	
		ACID PLANT PHYSIOL. 118 (4), 1536 (1998))	
6013	812	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE, 3'	Transferases
		PARTIAL SIMILAR TO	
		PHOSPHORIBOSYLANTHRANILATE TRANSFERASE	
		GB:CAA16616 [ARABIDOPSIS THALIANA][PUTATIVE]	
6049	813	DNA-3-METHYLADENINE GLYCOSYLASE I SIMILAR TO	Glycosylase
		PUTATIVE DNA-3-METHYLADENINE GLYCOSYLASE I	
		(TAG I) GB:P05100 [ESCHERICHIA COLI][PUTATIVE]	
6063	814	3-METHYLADENINE DNA GLYCOSYLASE, PUTATIVE	Glycosylase
		SIMILAR TO GI:1669515 FROM	
		[SCHIZOSACCHAROMYCES POMBE] (GENE 177 (1-2),	
		229-235 (1996))	
6067	815	GSH-DEPENDENT DEHYDROASCORBATE REDUCTASE	Reductase
]	!	1, PUTATIVE SIMILAR TO GI:6939839 FROM [ORYZA	
		SATIVA]	
6068	816	NADPH OXIDOREDUCTASE, PUTATIVE SIMILAR TO	Reductase
		GI:1708420 FROM [ARABIDOPSIS THALIANA] (J. BIOL.	
		CHEM. 270 (44), 26224-26231 (1995))	

6069	817	NADPH OXIDOREDUCTASE, PUTATIVE SIMILAR TO	Reductase
		GI:1708420 FROM [ARABIDOPSIS THALIANA] (J. BIOL	
		CHEM. 270 (44), 26224-26231 (1995))	
6070	818	NADPH OXIDOREDUCTASE, PUTATIVE SIMILAR TO	Reductase
		GI:1708420 FROM [ARABIDOPSIS THALIANA] (J. BIOL.	
		CHEM. 270 (44), 26224-26231 (1995))	
6084	819	CYTOKININ OXIDASE, PUTATIVE SIMILAR TO	Oxidase
		GB:CAA77151 FROM [ZEA MAYS] (PLANT J. 17 (6), 615-	
		626 (1999))	
6085	820	PROTEASE, PUTATIVE SIMILAR TO SP:P36774 FROM	Protease
		[MYXOCOCCUS XANTHUS]	
6101	821	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE GB:AAB36558	
		GI:1684913 FROM [IPOMOEA NIL]	
6114	822	PECTINESTERASE, PUTATIVE SIMILAR TO	Esterase
		PECTINESTERASE GB:CAB08077 GI:1944575 FROM	
		[LYCOPERSICON ESCULENTUM]	
6117	823	RECEPTOR KINASE (CLVI) IDENTICAL TO RECEPTOR	1
		KINASE (CLV1) GB:AAB58929 GI:2160756 [ARABIDOPSIS	
		THALIANA]	
6139	824	RNA PSEUDOURIDYLATE SYNTHASE, PUTATIVE	Synthase
		CONTAINS PFAM PROFILE: PF00849 RNA	
		PSEUDOURIDYLATE SYNTHASE	
6143	825	S-ADENOSYL-METHIONINE-STEROL-C-	Transferases
		METHYLTRANSFERASE, PUTATIVE ALMOST	
		IDENTICAL TO S-ADENOSYL-METHIONINE-STEROL-C-	
		METHYLTRANSFERASE GI:2246456 FROM	
6146	926	[ARABIDOPSIS THALIANA]	G4'
6146	826	PSEUDOURIDYLATE SYNTHASE 2 GI:1403092 FROM	Synthase
		[SACCHAROMYCES CEREVISIAE][HYPOTHETICAL	
(147	927	PROTEIN CONTAINS SIMILARITY TO] ALPHA-AMYLASE, PUTATIVE SIMILAR TO ALPHA-	Ghranulasa
6147	821	i e e e e e e e e e e e e e e e e e e e	Giycosyiase
6150	920	AMYLASE GI:7532799 FROM [ MALUS DOMESTICA] PECTINESTERASE, PUTATIVE SIMILAR TO	Esterase
6150	828	<b>'</b>	
		PECTINESTERASE GI:1944574 FROM [LYCOPERSICON	
		ESCULENTUM]	

6170	829	PROTEIN KINASE SIMILAR TO PROTEIN KINASE	Kinase, Protein
		(APK1A); CONTAINS PFAM PROFILE: PF00069	1
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]	
6176	830	PHOSPHATE TRANSPORTER SIMILAR TO PHOSPHATE	Transporter
		TRANSPORTER GB:BAA20522 [CATHARANTHUS	• •
		ROSEUS][PUTATIVE]	
6180	831	CINNAMOYL-COA REDUCTASE SIMILAR TO	Reductase
		CINNAMOYL-COA REDUCTASE GB:CAA56103	
		  [EUCALYPTUS GUNNII][PUTATIVE]	
6188	832	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE-	Kinase
		DEPENDENT PHOSPHOFRUCTOKINASE ALPHA	
	<b>[</b>	SUBUNIT SIMILAR TO PYROPHOSPHATE-DEPENDENT	
		PHOSPHOFRUCTOKINASE ALPHA SUBUNIT	
		GB:AAD30596 [ARABIDOPSIS THALIANA][PUTATIVE]	
6201	833	12-OXOPHYTODIENOATE REDUCTASE (OPR1)	Reductase
		IDENTICAL TO 12-OXOPHYTODIENOATE REDUCTASE	
ļ		OPR1 GB:AAC78440 [ARABIDOPSIS THALIANA]	
6202	834	12-OXOPHYTODIENOATE REDUCTASE (OPR2)	Reductase
		IDENTICAL TO 12-OXOPHYTODIENOATE REDUCTASE	
		OPR2 GB:AAC78441 [ARABIDOPSIS THALIANA]	
6212	835	CATECHOL O-METHYLTRANSFERASE SIMILAR TO	Transferases
		CATECHOL O-METHYLTRANSFERASE GB:CAA55358	
		[VANILLA PLANIFOLIA][PUTATIVE]	
6232	836	URIDYLYL TRANSFERASE GI:4406764 FROM	Transferases
		[ARABIDOPSIS THALIANA]DEACETYLVINDOLINE 4-0-	
		ACETYLTRANSFERASE, GB:AAC9931; UNKNOWN	
		PROTEIN SIMILAR TO UNKNOWN PROTEIN SIMILAR	
		TO PUTATIVE	
6238	837	CARBOXYPHOSPHONOENOLPYRUVATE MUTASE,	Mutase
		PUTATIVE SIMILAR TO	
		CARBOXYPHOSPHONOENOLPYRUVATE MUTASE	
		GI:47149 FROM [STREPTOMYCES HYGROSCOPICUS]	
6242	838	PEROXIDASE, PUTATIVE SIMILAR TO PEROXIDASE	Oxidase
		GI:1161565 FROM [LYCOPERSICON ESCULENTUM]	
6243	839	AUXIN TRANSPORT PROTEIN EIR1, PUTATIVE SIMILAR	Transporter
		TO AUXIN TRANSPORT PROTEIN EIR1 GI:3377507 FROM	
		[ARABIDOPSIS THALIANA]	

6259	840	RECEPTOR-LIKE PROTEIN KINASE GI:7529754 FROM Kinase, Protein
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO]
60.5	041	
6264	841	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE, Oxidase
	1	PUTATIVE SIMILAR TO 1-AMINOCYCLOPROPANE-1-
		CARBOXYLATE OXIDASE GI:3386565 FROM [SORGHUM]
		BICOLOR]
6269	842	AMINO ACID CARRIER, PUTATIVE SIMILAR TO AMINO Transporter
		ACID CARRIER GI:3293031 FROM [RICINUS COMMUNIS]
6274	843	LYSOPHOSPHOLIPASE ISOLOG, PUTATIVE SIMILAR TO Lipase
		LYSOPHOSPHOLIPASE ISOLOG GI:1931639 FROM
		[ARABIDOPSIS THALIANA]
6279	844	ASPARTIC PROTEASE, NUCELLIN-LIKE PROTEIN Protease
		SIMILAR TO NUCELLIN GB:AAB96882 [HORDEUM
		VULGARE] (NUCELLIN: SIMILAR TO ASPARTIC
		PROTEASE AND ITS SPECIFIC EXPRESSION IN
		NUCELLAR CELLS DURING DEGENERATION)
6280	845	ASCORBATE PEROXIDASE IDENTICAL TO Oxidase
		THYLAKOID-BOUND ASCORBATE PEROXIDASE
		GB:CAA67426 [ARABIDOPSIS THALIANA] THYLAKOID-
		BOUND
6283	846	CAFFEIC ACID 3-O-METHYLTRANSFERASE SIMILAR Transferases
		TO CAFFEIC ACID 3-0-METHYLTRANSFERASE
		GB:O23760 [CLARKIA BREWERI][PUTATIVE]
6284	847	CAFFEIC ACID 3-O-METHYLTRANSFERASE SIMILAR Transferases
		TO CAFFEIC ACID 3-O-METHYLTRANSFERASE
		GB:O23760 [CLARKIA BREWERI][PUTATIVE]
6299	848	AUX1-LIKE PERMEASE SIMILAR TO AUX1 Transporter
		(REGULATOR OF ROOT GRAVITROPISM, PUTATIVE
1		PERMEASE) GB:CAA67308 [ARABIDOPSIS
		THALIANA][PUTATIVE]
6308	849	ENDO-1,3-BETA-GLUCANASE SIMILAR TO ENDO-1,3-Glycosylase
		BETA-GLUCANASE GB:BAA21110 [GOSSYPIUM]
	] ]	HIRSUTUM][PUTATIVE]
6309		ENDO-1,3-BETA-GLUCANASE SIMILAR TO ENDO-1,3-Glycosylase
0309		
		BETA-GLUCANASE GB:BAA21110 [GOSSYPIUM]
	· ·	HIRSUTUM][PUTATIVE]

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6312	851	REVERSE TRANSCRIPTASE GI:976278 FROM Transcriptase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		CONTAINS SIMILARITY TO]
6321	852	NADH DEHYDROGENASE SUBUNIT 1[HYPOTHETICAL Dehydrogenase
j		PROTEIN CONTAINS SIMILARITY TO]
6335	853	PHOSPHOGLYCERATE MUTASE, PUTATIVE SIMILAR Mutase
		TO PHOSPHOGLYCERATE MUTASE GI:8979237 FROM
		[CHLAMYDIA MURIDARUM]
6336	854	XYLOSIDASE, PUTATIVE SIMILAR TO XYLOSIDASE Glycosylase
ļ		GI:2102655 FROM [ASPERGILLUS NIGER]
6339	855	TREHALOSE-6-PHOSPHATE PHOSPHATASE (ATTPPB) Phosphatase
	Ì	IDENTICAL TO TREHALOSE-6-PHOSPHATE
		PHOSPHATASE (ATTPPB) GB:AF007779 [ARABIDOPSIS
		THALIANA] (TREHALOSE SYNTHESIS, A COMMON
		DISACCHARIDE THAT APPEARS TO PLAY A MAJOR
		ROLE IN DESICCATION TOLERANCE, PLANT J 1998
		MAR;13(5):673-83)
6344	856	TRANSPORTER CONTAINS PFAM PROFILE: PF00083 Transporter
		SUGAR (AND OTHER) TRANSPORTER[PUTATIVE]
6351	857	PROTEIN PHOSPHATASE 2C SIMILAR TO PROTEIN Phosphatase
		PHOSPHATASE 2C GB:CAA72341 [MEDICAGO SATIVA];
	<b> </b> 	CONTAINS PFAM PROFILE: PF00481 PROTEIN
		PHOSPHATASE 2C[PUTATIVE]
6355	858	METHYLTRANSFERASE GB:AAD17428 [ARABIDOPSIS Transferases
		THALIANA]; UNKNOWN PROTEIN, 5' PARTIAL SIMILAR
6257	950	TO PUTATIVE
6357	839	UDP-GLUCOSE GLUCOSYLTRANSFERASE, PUTATIVE Transferases
		SIMILAR TO UDP-GLUCOSE GLUCOSYLTRANSFERASE
6362	960	GI:3928543 FROM [ARABIDOPSIS THALIANA]
0302	800	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO Transferases GLUTATHIONE TRANSFERASE GI:2853219 FROM
		[CARICA PAPAYA]
6363	961	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO Transferases
0303	601	GLUTATHIONE TRANSFERASE, POTATIVE SIMILAR TO Transierases  GLUTATHIONE TRANSFERASE GI:2853219 FROM
		[CARICA PAPAYA]
		[CARCATAFATA]

PCT/EP01/09892

6364	862	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO Transf	erases
		GLUTATHIONE TRANSFERASE GI:2853219 FROM	
		[CARICA PAPAYA]	
6365	863	GLUTATHIONE S-TRANSFERASE, PUTATIVE SIMILAR Transf	erases
		TO 2,4-D INDUCIBLE GLUTATHIONE S-TRANSFERASE	
		GI:2920666 FROM [GLYCINE MAX]	
6366	864	GLUTATHIONE TRANSFERASE, PUTATIVE SIMILAR TO Transfe	erases
		GLUTATHIONE TRANSFERASE GI:2853219 FROM	
		[CARICA PAPAYA]	
6368	865	EXOPOLYGALACTURONASE, PUTATIVE SIMILAR TO Glycos	vlase
		EXOPOLYGALACTURONASE GI:311962 FROM	J
		[ARABIDOPSIS THALIANA]	İ
6372	866	GIBBERELLIN 2- OXIDASE IDENTICAL TO Oxidas	e
ļ		GIBBERELLIN 2- OXIDASE GI:4678366 FROM	
		[ARABIDOPSIS THALIANA]	
6376	867	BETA-AMYRIN SYNTHASE, PUTATIVE SIMILAR TO Synthas	se
		BETA-AMYRIN SYNTHASE GI:3688600 FROM [PANAX]	
		GINSENG]	
6380	868	BETA-1,3-GLUCANASE, C TERMINAL FRAGMENT Glycos	ylase
	!	GI:2281103 FROM [ARABIDOPSIS	
1		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
6381	869	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TO Kinase,	, Protein
		RECEPTOR PROTEIN KINASE GI:9280288 FROM	
		[ARABIDOPSIS THALIANA]	
6383	870	FLAVANONE 3-HYDROXYLASE, PUTATIVE SIMILAR Hydrox	ylase
		TO FLAVANONE 3-HYDROXYLASE GI:727410 FROM	
		[PERSEA AMERICANA]	.
6420	871	INORGANIC PYROPHOSPHATASE IDENTICAL TO Phosph	atase
		VACUOLAR-TYPE H+-TRANSLOCATING INORGANIC	ļ
		PYROPHOSPHATASE GI:6901678 FROM [ARABIDOPSIS	
		THALIANA]	
6426	872	LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN Kinase,	Protein
		KINASE, PUTATIVE SIMILAR TO LEUCINE-RICH	Ì
		REPEAT TRANSMEMBRANE PROTEIN KINASE 2	
		GI:3360291 FROM [ZEA MAYS]	
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6465	873	NA+/H+ ANTIPORTER GI:4558666 FROM [ARABIDOPSIS]T	ransporter
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
6471	874	ENT-KAURENE SYNTHASE, PUTATIVE SIMILAR TOS	ynthase
		ENT-KAURENE SYNTHASE GI:3056725 FROM	
		[ARABIDOPSIS THALIANA]	
6473	875	BETA-1,3-GLUCANASE-LIKE PROTEIN GI:9758115 FROM G	lycosylase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN	
]		SIMILAR TO]	
6475	876	2-DEHYDRO-3-DEOXYPHOSPHOOCTONATE	ldolase
		ALDOLASE, PUTATIVE SIMILAR TO 2-DEHYDRO-3-	
		DEOXYPHOSPHOOCTONATE ALDOLASE GI:1683630	
		FROM [MANNHEIMIA HAEMOLYTICA]	ļ
6486	877	NA+/H+ ANTIPORTER GI:1655701 FROM [XENOPUS T	ransporter
		LAEVIS][HYPOTHETICAL PROTEIN CONTAINS	
		SIMILARITY TO]	•
6487	878	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TOK	inase, Protein
		RECEPTOR PROTEIN KINASE GI:1389566 FROM	
		[ARABIDOPSIS THALIANA]	
6488	879	PROTEIN PHOSPHATASE-2C, PUTATIVE SIMILAR TO PI	nosphatase
		PROTEIN PHOSPHATASE-2C GI:3643088 FROM	
		[MESEMBRYANTHEMUM CRYSTALLINUM]	
6492	880	WALL-ASSOCIATED KINASE 4, PUTATIVE SIMILAR TO K	inase, Protein
		WALL-ASSOCIATED KINASE 4 GI:3355308 FROM	
		[ARABIDOPSIS THALIANA]	
6493	881	WALL-ASSOCIATED KINASE 2, PUTATIVE SIMILAR TO K	inase, Protein
		WALL-ASSOCIATED KINASE 2 GI:4826399 FROM	
		[ARABIDOPSIS THALIANA]	
6532	882	RECEPTOR PROTEIN KINASE, PUTATIVE SIMILAR TOKE	inase, Protein
		RECEPTOR PROTEIN KINASE GI:1389566 FROM	į
5500		[ARABIDOPSIS THALIANA]	
6538	883		lycosylase
		POLYGALACTURONASE GI:7381227 FROM	
CEAN	004	[LYCOPERSICON ESCULENTUM]	h vo av le
6541	884	·	lycosylase
	: 	POLYGALACTURONASE GI:7381227 FROM	
		[LYCOPERSICON ESCULENTUM]	

PCT/EP01/09892

6551	885	ALPHA/BETA HYDROLASE FOLD	Hydrolase
6554	886	SULFATE TRANSPORTER CONTAINS PFAM PROFILE:	
		PF00916 SULFATE TRANSPORTER FAMILY[PUTATIVE]	1
6555	887	IRON/ASCORBATE OXIDOREDUCTASE	Reductase
		FAMILY[PUTATIVE]	ļ
6556	888	GIBBERELLIN 3 BETA-HYDROXYLASE SIMILAR TO	Hydroxylase
		GIBBERELLIN 3 BETA-HYDROXYLASE GB:AAC83647	
		[ARABIDOPSIS THALIANA][PUTATIVE]	]
6557	889	GIBBERELLIN 3 BETA-HYDROXYLASE IDENTICAL TO	Hydroxylase
		GIBBERELLIN 3 BETA-HYDROXYLASE GB:AAC83647	
		[ARABIDOPSIS THALIANA]	
6559	890	ASPARTATE AMINOTRANSFERASE SIMILAR TO	Transferases
		ASPARTATE AMINOTRANSFERASE B GB:Q06191	
		[SINORHIZOBIUM MELILOTI][PUTATIVE]	
6573	891	AMINO ACID TRANSPORTER PROTEIN DOMAIN	Transporter
6586	892	PROTEIN KINASE SIMILAR TO LEUCINE-RICH REPEAT	Kinase, Protein
		TRANSMEMBRANE PROTEIN KINASE I GB:AAC27894	
		FROM [ZEA MAYS][PUTATIVE]	
6604	893	CINNAMOYL COA REDUCTASE, PUTATIVE SIMILAR TO	Reductase
		CINNAMOYL COA REDUCTASE GB:AAF43141	
		GI:7239228 FROM [POPULUS TREMULOIDES]	
6609	894	PROTEIN KINASE, PUTATIVE SIMILAR TO RECEPTOR-	Kinase, Protein
		LIKE SERINE/THREONINE KINASE GB:AAC50043	
		GI:2465923 FROM [ARABIDOPSIS THALIANA]	
6633	895	į i	Transporter
		GP:AF019637[PUTATIVE]	
6658	896		Transporter
		GP 3785972 AC005560[PUTATIVE]	
6663	897	PROTEIN TRANSPORT PROTEIN SEC12P IDENTICAL TO	Transporter
		GB:M95796, CONTAINS A WD-40 REPEAT	
		DOMAIN[PUTATIVE]	
6679		BETA-1,3-GLUCANASE [PUTATIVE]	Glycosylase
6699	899	HISTIDINE KINASE [PUTATIVE]	Kinase, Protein
6701	900	XYLOGLUCAN-SPECIFIC GLUCANASE IDENTICAL TO	Glycosylase
}		GB:D63509[PUTATIVE]	

(704	001	DUDDIE ACID DISCOULATAGE CONTADIO ACTUALIO	In .
6704	901	PURPLE ACID PHOSPHATASE CONTAINS METALLO	- Esterase
570.5	000	PHOSPHOESTERASE MOTIF (PS50185)[PUTATIVE]	
6705	902	PURPLE ACID PHOSPHATASE CONTAINS METALLO	Esterase
		PHOSPHOESTERASE MOTIF (PS50185)[PUTATIVE]	<u> </u>
6711	903		Kinase, Protein
		BRASSINOSTEROID INSENSITIVE PROTEIN[PUTATIVE]	}
6714	904	NA+/H+ ANTIPORTER [PUTATIVE]	Transporter
6716	905	GLUTAMATE DECARBOXYLASE [PUTATIVE]	Decarboxylase
6717	906	GLUTAMATE DECARBOXYLASE [PUTATIVE]	Decarboxylase
6737	907	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
6752	908	CINNAMOYL-COA REDUCTASE [PUTATIVE]	Reductase
6759	909	DNA POLYMERASE III GAMMA SUBUNIT [SIMILAR TO]	Polymerase
6763	910	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
6770	911	TRANSPORT PROTEIN [PUTATIVE]	Transporter
6775	912	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
6781	913	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
6782	914	PECTATE LYASE [PUTATIVE]	Lyase
6788	915	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
6801	916	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
6806	917	RIBONUCLEASE, RNS1 IDENTICAL TO GB:U05206;	Nuclease
		CONTAINS A RIBONUCLEASE T2 FAMILY HISTIDINE	
		ACTIVE SITE SIGNATURE (PDOC00459)[PUTATIVE]	
6844	918	HORMONE-RECEPTOR DOMAIN (PROSITE: QDOC50227)	Receptor
6860	919	ESTERASE (CONTAINS AN	Esterase
		ESTERASE/LIPASE/THIOESTERASE ACTIVE SITE	:
		SERINE DOMAIN (PROSITE: PS50187); RELATED TO	
		PLANT SENSITIVE RESPONSE PROTEINS[PUTATIVE]	
6875	920	URIDYLYL TRANSFERASE [PUTATIVE]	Transferases
6877	921	STEROID SULFOTRANSFERASE [PUTATIVE]	Transferases
6878	922	STEROID SULFOTRANSFERASE [PUTATIVE]	Transferases
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6890	923	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APO	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
6892	924	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APO	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
6894	925	IRON TRANSPORTER ROOT [PUTATIVE]	Transporter
6908	926	RIBONUCLEASE E [PUTATIVE]	Nuclease
6910	927	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
6920	928	INDOLE-3-GLYCEROL PHOSPHATE SYNTHASE	Synthase
		[PUTATIVE]	
6922	929	NON-LTR RETROLELEMENT REVERSE	Transcriptase
		TRANSCRIPTASE [PUTATIVE]	
6933	930	PROTEIN PHOSPHATASE [PUTATIVE]	Phosphatase
6935	931	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
6936	932	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
6946	933	FERREDOXIN-THIOREDOXIN REDUCTASE [PUTATIVE]	Reductase
6952	934	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
6980	935	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
6986	936	PHOSPHORIBOSYLAMINOIMIDAZOLE CARBOXYLASE	Carboxylase
		[PUTATIVE]	
7023	937	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
7038	938	BETA-1,3-GLUCANASE [PUTATIVE]	Glycosylase
7050	939	SERINE PROTEASE [PUTATIVE]	Protease
7057	940	ENOYL-ACP REDUCTASE (ENR-A) IDENTICAL TO	Reductase
		GB:Y13860	
7063	941	12-OXOPHYTODIENOATE-10,11-REDUCTASE	Reductase
7110	942	ENDOXYLOGLUCAN GLYCOSYLTRANSFERASE	Transferases
		IDENTICAL TO GB:D16454[PUTATIVE]	
7115	943	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
7123	944	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein

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7125	945	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
7126	946	,	Protease
		PARTIAL[PUTATIVE]	-
7133	947	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
7168		ABC TRANSPORTER [PUTATIVE]	Transporter
7262	949	MONOGALACTOSYLDIACYLGLYCEROL SYNTHASE	Synthase
		[PUTATIVE]	
7263	950	REVERSE TRANSCRIPTASE TA1-1;	Transcriptase
		GB:AAD17398[HYPOTHETICAL PROTEIN SIMILAR TO]	
7295	951	C-I-TETRAHYDROFOLATE SYNTHASE [PUTATIVE]	Synthase
7331	952	N-ACETYLGLUCOSAMINYLTRANSFERASE	Transferases
		[PUTATIVE]	
7352	953	NA/H ANTIPORTER [PUTATIVE]	Transporter
7365	954	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7366	955	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7367	956	ASPARTATE AMINOTRANSFERASE [PUTATIVE]	Transferases
7403	957	PROLINE IMINOPEPTIDASE IDENTICAL TO	Protease
		GP:1710151:U72711	
7404	958	PROTEIN PHOSPHATASE 2C [PUTATIVE]	Phosphatase
7436	959	ENDOXYLOGLUCAN GLYCOSYLTRANSFERASE	Transferases
]		[PUTATIVE]	
7441	960	SUCROSE-PROTON SYMPORTER [PUTATIVE]	Transporter
7443	961	1,4-BETA-XYLAN ENDOHYDROLASE	Hydrolase
7446	962	VACUOLAR SORTING RECEPTOR-LIKE PROTEIN	Receptor
7448	963	VACUOLAR SORTING RECEPTOR-LIKE PROTEIN	Receptor
7465	964	STEROID SULFOTRANSFERASE [PUTATIVE]	Transferases
7500	965	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7504	966	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
7509	967	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	

7516	968	POLYGALACTURONASE [PUTATIVE]	Glycosylase
7517	969	POLYGALACTURONASE [PUTATIVE]	Glycosylase
7518	970	POLYGALACTURONASE [PUTATIVE]	Glycosylase
7519	971	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
7520	972	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
7522	973	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
7533	974	FERREDOXINNITRITE REDUCTASE	Reductase
7543	975	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
7546	976	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
7550	977	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE CYCLOPHILIN-TYPE [PUTATIVE]	Isomerase
7577	978	NON-LTR RETROLELEMENT REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
7588	979	BETA-1,3-ENDOGLUCANASE, 5' PARTIAL [PUTATIVE]	Glycosylase
7589	980	LRR RECEPTOR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7592	981	BETA-KETOACYL-COA SYNTHASE [PUTATIVE]	Synthase
7604	982	PURPLE ACID PHOSPHATASE PRECURSOR [PUTATIVE]	Phosphatase
7612	983	ARGININE DECARBOXYLASE IDENTICAL TO GP:1590814:U52851	Decarboxylase
7623	984	PROTEIN KINASE CONTAINS A PROTEIN KINASE DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Kinase, Protein
7636	985	PROTEIN KINASE CONTAINS A PROTEIN KINASE DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Kinase, Protein
7640	986	GLUCONOKINASE [PUTATIVE]	Kinase
7651	987	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
7659		TETRACYCLINE TRANSPORTER PROTEIN [PUTATIVE]	Transporter
7660	989	TETRACYCLINE TRANSPORTER PROTEIN [PUTATIVE]	Transporter
7661	990	TETRACYCLINE TRANSPORTER-LIKE PROTEIN, 3'	Transporter
		PARTIAL IDENTICAL TO EGAD 110957 119053	
7664	991	PROTEIN KINASE DOMAIN	Kinase, Protein

7685	992	POLY(A) POLYMERASE [PUTATIVE]	Polymerase
7688	993	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
7691	994	SERINE ACETYLTRANSFERASE (GB:AF112303)	Transferases
7708	995	SERINE PEPTIDASE RELATED TO THYMUS-SPECIFIC	Protease
		FROM HOMO SAPIENS	
7714	996	PURPLE ACID PHOSPHATASE PRECURSOR [PUTATIVE]	Phosphatase
7715	997	PEROXIDASE [PUTATIVE]	Oxidase
7716	998	PEROXIDASE [PUTATIVE]	Oxidase
7720	999	AAA-TYPE ATPASE [PUTATIVE]	ATPase
7724		INORGANIC PYROPHOSPHATASE [PUTATIVE]	Phosphatase
7746	1001	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
7752	1002	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
	·	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
7755	1003	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
7756	1004	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
7761	1005	GERANYLGERANYL PYROPHOSPHATE SYNTHASE	Synthase
		[PUTATIVE]	
7763	1006	GERANYLGERANYL PYROPHOSPHATE SYNTHASE	Synthase
		CONTAINS GB:L22347[PUTATIVE]	
7779	1007	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE	Glycosylase
		[PUTATIVE]	
7789		PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
7798	1009	PEROXIDASE (ATP22A) IDENTICAL TO GB:Y08781	Oxidase
7801	1010	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
7805	1011	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	ļ
7806	1012	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	

7807	1012	ANTHRANILATE	Transferases
7807	1013		- I ransierases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE	
		[PUTATIVE]	
7810	1014	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
7813	1015	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7815	1016	PECTINESTERASE [PUTATIVE]	Esterase
7817	1017	SERINE PROTEASE [PUTATIVE]	Protease
7819	1018	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7821	1019	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
7823	1020	PROTEIN KINASE, 5'PARTIAL [PUTATIVE]	Kinase, Protein
7844	1021	BETA-1,3-ENDOGLUCANASE [PUTATIVE]	Glycosylase
7850	1022	CYTOKININ OXIDASE [PUTATIVE]	Oxidase
7855	1023	ALPHA/BETA HYDROLASE FOLD	Hydrolase
		(PF00561).[PUTATIVE]; ESTERASE CONTAINS	
	1	ESTERASE/LIPASE/THIOESTERASE ACTIVE SITE	
		SERINE (PS50187)	
7857	1024	CYTIDINE DEAMINASE [PUTATIVE]	Deaminase
7859	1025	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
7860	1026	POTASSIUM/PROTON ANTIPORTER [PUTATIVE]	Transporter
7894	1027	N-ACETYL-GAMMA-GLUTAMYL-PHOSPHATE	Reductase
		REDUCTASE [PUTATIVE]	
7905	1028	PROTEIN PHOSPHATASE 2C [PUTATIVE]	Phosphatase
7930	1029	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Í
7962		PROTEIN PHOSPHATASE 2C [PUTATIVE]	Phosphatase
7969	1031	(1-4)-BETA-MANNAN ENDOHYDROLASE	Hydrolase
7970			Synthase
7985		SECRETORY CARRIER-ASSOCIATED MEMBRANE	
		PROTEIN [PUTATIVE]	
7986	1034	LRR RECEPTOR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8002	1035	MAJOR INTRINSIC (CHANNEL) PROTEIN [PUTATIVE]	Channel
8005	1036	AUX1-LIKE AMINO ACID PERMEASE	Transporter
		·	

		·	
8016	1037	SIGNAL SEQUENCE RECEPTOR, ALPHA SUBUNIT (SSR-	Receptor
		ALPHA) SAME AS GP: 1174448[PUTATIVE]	
8036	1038	XYLULOSE KINASE [PUTATIVE]	Kinase
8047	1039	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
	,	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8054	1040	DIHYDROFOLATE REDUCTASE, 3' PARTIAL	Reductase
		[PUTATIVE]	
8059	1041	PECTINESTERASE [PUTATIVE]	Esterase
8072	1042	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
•		[PUTATIVE]	
8084	1043	CINNAMYL-ALCOHOL DEHYDROGENASE [PUTATIVE]	Dehydrogenases
8089	1044	SHIKIMATE KINASE PRECURSOR [PUTATIVE]	Kinase
8113	1045	TREHALOSE-6-PHOSPHATE PHOSPHATASE	Phosphatase
		[PUTATIVE]	
8115	1046	BETA-HYDROXYACYL-ACP DEHYDRATASE	Dehydratase
		[PUTATIVE]	
8131	1047	PEROXIDASE [PUTATIVE]	Oxidase
8133	1048	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
8134	1049	GTP CYCLOHYDROLASE [PUTATIVE]	Hydrolase
8137	1050	PYROPHOSPHATEFRUCTOSE-6-PHOSPHATE 1-	Transferases
		PHOSPHOTRANSFERASE [PUTATIVE]	<u>.</u>
8147	1051	ANTHOCYANIDIN-3-GLUCOSIDE	Transferases
		RHAMNOSYLTRANSFERASE, 3' PARTIAL	
8153	1052	D-AMINO ACID DEHYDROGENASE [PUTATIVE]	Dehydrogenases
8179	1053	AMINO ACID ACETYLTRANSFERASE [PUTATIVE]	Transferases
8181	1054	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8184	1055	SERINE CARBOXYPEPTIDASE I, PSEUDOGENE SIMILAR	Protease
		TO SERINE CARBOXYPEPTIDASE I[PUTATIVE]	
8207	1056	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8208	1057	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8210	1058	VETISPIRADIENE SYNTHASE [PUTATIVE]	Synthase

8212	1059	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8213	1060	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8217	1061	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8232	1062	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8238	1063	ANTHRANILATE N-	Transferases
	:	HYDROXYCINNAMOYL/BENZOYLTRANSFERASE	
	:	[PUTATIVE]	
8241	1064	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
	!	PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
8242	1065	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8243	1066	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8244	1067	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8245	1068	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8246	1069	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8247	1070	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8248	1071	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8249	1072	ACETONE-CYANOHYDRIN LYASE [PUTATIVE]	Lyase
8250	1073	PECTINESTERASE [PUTATIVE]	Esterase
8263	1074	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8266	1075	GERANYLGERANYL PYROPHOSPHATE SYNTHASE	Synthase
		IDENTICAL TO GB:U44876; SEQUENCE DISAGREES AT	
		N-TERMINUS, SEQUENCE SUBMITTED HAS BEEN	
		CONFIRMED FROM THREE ELECTROPHEROGRAMS.	
8274	1076	POLYGALACTURONASE [PUTATIVE]	Glycosylase
8275	1077	CINNAMOYL COA REDUCTASE [PUTATIVE]	Reductase
8279	1078	LRR RECEPTOR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8282	1079	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL	
8296	1080	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8304	1081	LIMONENE CYCLASE [PUTATIVE]	Cyclase
8334	1082	H+ ATPASE, PLASMA MEMBRANE, 3' PARTIAL	ATPase
		[PUTATIVE]	
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PCT/EP01/09892

6226	1002	I IDACE/UVDBOLACE CDCI	)r :
8338	1083	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
8343	l	CYCLIC NUCLEOTIDE-REGULATED ION CHANNEL	Channel
		PROTEIN [PUTATIVE]	
8345	1085	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8353	1086	LIGAND-GATED ION CHANNEL PROTEIN [PUTATIVE]	Channel
8354	1087	LIGAND-GATED ION CHANNEL PROTEIN [PUTATIVE]	Channel
8358	1088	PEROXIDASE [PUTATIVE]	Oxidase
8399	1089	PROTEIN KINASE DOMAIN	Kinase, Protein
8420	1090	DIOXYGENASE [PUTATIVE]	Oxygenases
8429	1091	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
		[PUTATIVE]	
8455	1092	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8498	1093	BETA-KETOACYL-COA SYNTHASE (FIDDLEHEAD)	Synthase
		IDENTICAL TO GB:AJ010713; CONTAINS A CHALCONE	
		AND STILBENE SYNTHASE ACTIVE SITE (PF00195)	
8502	1094	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8506	1095	RECEPTOR-LIKE PROTEIN KINASE, ERECTA	Kinase, Protein
		IDENTICAL TO GB:U47029 AND GB:D83257; CONTAINS	
		A PROTEIN KINASE DOMAIN PROFILE (PDOC00100);	Ī
		CONTAINS LRR LEUCINE RICH REPEAT	
		DOMAINS[PUTATIVE]	
8509	1096	MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE]	Transporter
8517	1097	PECTINESTERASE [PUTATIVE]	Esterase
8518	1098	PECTINESTERASE [PUTATIVE]	Esterase
8521	1099	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8526	1100	PHEROMONE RECEPTOR AR781, SIMILAR TO YEAST;	Receptor
		IDENTICAL TO GB:D88743, CORRECTED A FRAMESHIFT	
		FOUND IN THE ORIGINAL RECORD (AT 69530 BP),	į
		SEQUENCE SUBMITTED HAS BEEN VERIFIED FROM 10	
		SEQUENCE ELECTROPHEROGRAMS. THE	
]		TRANSLATION NOW STARTS FROM AN UPSTREAM	
		ATG.	
<u> </u>			

8528	1101	HEME OXYGENASE 2 (HO2)	Oxygenases
8533	1102	BETA-1,3-GLUCANASE [PUTATIVE]	Glycosylase
8535	1103	POLYGALACTURONASE [PUTATIVE]	Glycosylase
8537	1104	BETA-KETOACYL-COA SYNTHASE [PUTATIVE]	Synthase
8540	1105	·	Oxygenases
		GB:AF132475; ANNOTATION UPDATED PER SETH J.	
	1106	DAVIS AT UNIVERSITY OF WISCONSIN-MADISON	
8546	1106	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8560		PHOSPHOLIPASE C [PUTATIVE]	Lipase
8563	1108	BILE ACID TRANSPORTER, NA+ DEPENDENT ILEAL	Transporter
		[PUTATIVE]	
8579	1109	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8609	1110	LIPASE [PUTATIVE]	Lipase
8611	1111	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
8624	1112	BETA-1,3-GLUCANASE [PUTATIVE]	Glycosylase
8630	1113	FLAVONOL SULFOTRANSFERASE [PUTATIVE]	Transferases
8655	1114	PREPHENATE DEHYDRATASE/CHORISMATE MUTASE	Dehydratase
		[PUTATIVE]	
8658	1115	DTDP-GLUCOSE 4-6-DEHYDRATASE [PUTATIVE]	Dehydratase
8659	1116	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		[PUTATIVE]	
8680	1117	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8693	1118	CARBONIC ANHYDRASE [PUTATIVE]	Anhydrase
8697	1119	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8698	1120	CYCLIC NUCLEOTIDE AND CALMODULIN-	Channel
		REGULATED ION CHANNEL PROTEIN [PUTATIVE]	
8719	1121	PHEROMONE RECEPTOR DEFICIENT MUTANT	Receptor
		[SIMILAR TO]	!
8758	1122	PARA-AMINOBENZOATE SYNTHASE AND GLUTAMINE	Synthase
		AMIDOTRANSFERASE, A BIFUNCTIONAL ENZYME	
		[PUTATIVE]	
8760	1123	MEMBRANE CHANNEL PROTEIN [PUTATIVE]	Channel

25.50			1
8769	1124	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
8778	1125	LIGAND-GATED ION CHANNEL PROTEIN [PUTATIVE]	Channel
8779	1126	LIGAND-GATED ION CHANNEL PROTEIN [PUTATIVE]	Channel
8780	1127	LIGAND-GATED ION CHANNEL PROTEIN [PUTATIVE]	Channel
8784	1128	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8789	1129	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8792	1130	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
	-	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8793	1131	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8794	1132	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8800	1133	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8801	1134	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8803	1135	PROTEIN PHOSPHATASE 2C [PUTATIVE]	Phosphatase
8807	1136	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
8809	1137	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
8810	1138	GLUTATHIONE S-TRANSFERASE IDENTICAL TO	Transferases
		GB:X89216	
8811	1139	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
0010	1140	CLUTATIONE OF TRANSPERACE INSTATISES	TC C
8812	1140	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
8813	1141	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
8814	1142	GLUTATHIONE S-TRANSFERASE [PUTATIVE]	Transferases
0014	1142	GLUTATRIONE S-TRANSFERASE [FUTATIVE]	Transferases
8834	1143	ANTHRANILATE SYNTHASE, ALPHA SUBUNIT	Synthase
		IDENTICAL TO GB:M92354	
8836	1144	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8837	1145	MONOOXYGENASE [PUTATIVE]	Oxygenases
8838	1146	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8839	1147	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8840	1148	FLAVONOL 3-O-GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases

8864	1140	NADH DEHYDROGENASE (UBIQUINONE	Dahadaa
8004	1149		Dehydrogenase
		OXIDOREDUCTASE) [PUTATIVE]	
8872	1150	HIGH-AFFINITY POTASSIUM TRANSPORTER (ATKUP1)	Transporter
		IDENTICAL TO GB:AF029876	
8873	1151	FE(II) TRANSPORT PROTEIN [PUTATIVE]	Transporter
8879	1152	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8880	1153	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
8885	1154	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
8887	1155	NA/H ANTIPORTER [PUTATIVE]	Transporter
8892	1156	VACUOLAR SORTING RECEPTOR [PUTATIVE]	Receptor
8894	1157	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
8895	1158	PSEUDOURIDINE SYNTHASE [PUTATIVE]	Synthase
8907	1159	SIGNAL PEPTIDASE I [PUTATIVE]	Protease
8917	1160	LIPASE [PUTATIVE]	Lipase
8929	1161	TROPINONE REDUCTASE [PUTATIVE]	Reductase
8935	1162	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8936	1163	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8945	1164	DIOXYGENASE [PUTATIVE]	Oxygenases
8946	1165	DIOXYGENASE [PUTATIVE]	Oxygenases
8947	1166	GLUTATHIONE S-TRANSFERASE IDENTICAL TO	Transferases
]		GB:Y12295	]
8948	1167	GLUTATHIONE S-TRANSFERASE IDENTICAL TO	Transferases
		GB:D17673	
8955	1168	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
	ĺ	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
8972	1169	LIPASE [PUTATIVE]	Lipase
8987	1170	GLUTAMYL TRNA REDUCTASE [PUTATIVE]	Reductase
8998	1171	DELTA 9 DESATURASE ALMOST IDENTICAL (4 AA	Desaturases
		DIFF'T) TO GP:2970036	- Parties
9001	1172	FRUCTOKINASE [PUTATIVE]	Kinase
LL	!		

9016	1173	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APO	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
9017	1174	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APO	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
9031	1175	TRIACYLGLYCEROL LIPASE [PUTATIVE]	Lipase
9038	1176	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
9042	1177	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
9043	1178	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9044	1179	ACETOLACTATE SYNTHASE [PUTATIVE]	Synthase
9054	1180	NA+/H+ ANTIPORTER [PUTATIVE]	Transporter
9060	1181	GLUCAN SYNTHASE [PUTATIVE]	Synthase
9066	1182	ALANINE ACETYL TRANSFERASE [PUTATIVE]	Transferases
9067	1183	ALANINE ACETYL TRANSFERASE [PUTATIVE]	Transferases
9078	1184	HYDROLASE [PUTATIVE]	Hydrolase
9090	1185	FE(II) TRANSPORTER IDENTICAL TO	Transporter
		GB:AF033537[PUTATIVE]	
9092	1186	BETA-AMYLASE [PUTATIVE]	Glycosylase
9108	1187	O-GLCNAC TRANSFERASE [PUTATIVE]	Transferases
9116	1188	CELLULOSE SYNTHASE [PUTATIVE]	Synthase
9117	1189	CELLULOSE SYNTHASE [PUTATIVE]	Synthase
9124	1190	CELLULOSE SYNTHASE [PUTATIVE]	Synthase
9125	1191	CELLULOSE SYNTHASE [PUTATIVE]	Synthase
9140	1192	PURPLE ACID PHOSPHATASE PRECURSOR [PUTATIVE]	Phosphatase
9143	1193	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	1
9146	1194	PHOSPHATE TRANSPORTER [PUTATIVE]	Transporter
9173	1195	CELLULOSE SYNTHASE [PUTATIVE]	Synthase
9178	1196	POLYGALACTURONASE [PUTATIVE]	Glycosylase
9179	1197	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
9185	1198	FLAVIN-CONTAINING MONOOXYGENASE [PUTATIVE]	Oxygenases
9188	1199	TYROSINE-SPECIFIC TRANSPORT PROTEIN	Transporter
	·	[PUTATIVE]	
	-		

DOMAIN PROFILE (PDOC00100)[PUTATIVE]  9221 1201 CINNAMOYL-COA REDUCTASE HIGHLY SIMILAR TO Reduct F4P9.37[PUTATIVE]  9222 1202 CINNAMOYL-COA REDUCTASE HIGHLY SIMILAR TO Reduct F4P9.36[PUTATIVE]  9242 1203 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp  9257 1204 PEROXIDASE [PUTATIVE] Oxidas	
F4P9.37[PUTATIVE]  9222 1202 CINNAMOYL-COA REDUCTASE HIGHLY SIMILAR TO Reduct F4P9.36[PUTATIVE]  9242 1203 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp	
9222 1202 CINNAMOYL-COA REDUCTASE HIGHLY SIMILAR TO Reduct F4P9.36[PUTATIVE] 9242 1203 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp	ase
F4P9.36[PUTATIVE] 9242 1203 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp	ase
9242 1203 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp	
9257 1204 PEROXIDASE [PUTATIVE] Oxidas	orter
	е
9280 1205 PROTEIN KINASE CONTAINS A PROTEIN KINASE Kinase	, Protein
DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9283 1206 NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transc	riptase
[PUTATIVE]	
9306 1207 GIBBERELLIN 2-OXIDASE [PUTATIVE] Oxidas	е
9325 1208 PROTEIN PHOSPHATASE 2C [PUTATIVE] Phosph	atase
9344 1209 VACUOLAR SORTING RECEPTOR [PUTATIVE] Recept	or
9347 1210 PHOSPHATIDYLINOSITOL-GLYCAN SYNTHASE Syntha:	se
[PUTATIVE]	ŀ
9355 1211 POTASSIUM TRANSPORTER [PUTATIVE] Transp	orter
9385 1212 PEROXIDASE [PUTATIVE] Oxidas	e
. 9409 1213 RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE] Kinase	, Protein
9412 1214 GLUCOSYLTRANSFERASE [PUTATIVE] Transfe	erases
9413 1215 MONOOXYGENASE [PUTATIVE] Oxyger	nases
9428 1216 MITOCHONDRIAL CARRIER PROTEIN [PUTATIVE] Transp	orter
9469 1217 PHOSPHORIBOSYLFORMIMINO-5-AMINOIMIDAZOLE Isomera	ase
CARBOXAMIDE RIBOTIDE ISOMERASE IDENTICAL TO	
AB006139	
9500 1218 RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE] Kinase,	Protein
9502 1219 PROLINE TRANSPORTER [PUTATIVE] Transpo	orter
9509 1220 ASPARTYL PROTEASE FAMILY (PF00026)[PUTATIVE] Proteas	е
9511 1221 GIBERELLIN BETA-HYDROXYLASE CONTAINS Hydrox	ylase
SIMILARITIES TO GA BETA-20-HYDROXYLASE FROM	
TOBACCO (GB:3327245) AND TO ETHYLENE FORMING	
ENZYME FROM PICEA GLAUCA	
(GB:L42466)[PUTATIVE]	İ

9512	1222	PECTINESTERASE [PUTATIVE]	Esterase
9513	1223	PECTINESTERASE [PUTATIVE]	Esterase
9517	1224	GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
9518	1225	GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
9519		GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
9520	1227	GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
9521	1228	GLUCOSYL TRANSFERASE AN EST MATCHING THE 5'	
		END OF THIS GENE (GB:AA605508) WAS ORIGINALLY	
		DESCRIBED AS POLYADENYLATED (GB:AA006321)	
		AND IS PROBABLY TRANSCRIBED FROM THE	:
		OPPOSITE STRAND[PUTATIVE]	
9522		GLUCOSYL TRANSFERASE [PUTATIVE]	Transferases
9527	1230	GLUCAN SYNTHASE [PUTATIVE]	Synthase
9528	1231	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE	Glycosylase
		[PUTATIVE]	
9538	1232	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
9540	1233	RNA POLYMERASE SIGMA-70 FACTOR [PUTATIVE]	Polymerase
9546	1234	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
9554	1235	PEROXIDASE ATP2A [PUTATIVE]	Oxidase
9555	1236	VETISPIRADIENE SYNTHASE [PUTATIVE]	Synthase
9591	1237	GLUTAMATE/ORNITHINE ACETYLTRANSFERASE [PUTATIVE]	Transferases
9611		PHOSPHORIBOSYLAMINOIMIDAZOLE CARBOXYLASE [PUTATIVE]	Carboxylase
9612	1239	CER1-LIKE PROTEIN MAY BE INVOLVED IN WAX	Desaturases
		BIOSYNTHESIS; CONTAINS A SUR2-TYPE	
		HYDROXYLASE/DESATURASE CATALYTIC DOMAIN	
	j	(PS50242)	
9613	1240	RECEPTOR-LIKE PROTEIN KINASE SAME AS GB:X95909	Kinase, Protein
ł		(POLYMORPHISM EXISTS AT A GA REPEAT. WE	
1		FOUND 6 COPIES IN OUR SEQUENCE WHEREAS ONLY 5	
		COPIES EXIST IN GB:X95909)[PUTATIVE]	
<u>.</u> J.			

9645	1241	ALPHA-CARBOXYLTRANSFERASE [PUTATIVE]	Transferases
9651	1242	PEPTIDE/AMINO ACID TRANSPORTER [PUTATIVE]	Transporter
9658	1243	HIGH AFFINITY CA2+ ANTIPORTER IDENTICAL TO	Transporter
,		GB:U57411, EXCEPT A POSSIBLE FRAMESHIFT AT BASE	
		58008. SEQUENCE HAS BEEN CONFIRMED WITH 5	
		SEQUENCING READS.	
9665	1244	ANTHOCYANIDIN SYNTHASE [PUTATIVE]	Synthase
9669	1245	AMMONIUM TRANSPORTER [PUTATIVE]	Transporter
9678	1246	PEROXIDASE IDENTICAL TO M58381	Oxidase
9679	1247	PEROXIDASE	Oxidase
9700	1248	ACID PHOSPHATASE CONTAINS METALLO-	Esterase
		PHOSPHOESTERASE MOTIF (PS50185)[PUTATIVE]	
9734	1249	PHOSPHATE TRANSPORTER (ATPT2) IDENTICAL TO	Transporter
		GB:U62331	
9743	1250	PEROXIDASE [PUTATIVE]	Oxidase
9750	1251	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9757	1252	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	1
9758	1253	ABC TRANSPORTER [PUTATIVE]	Transporter
9766	1254	ADENYLATE KINASE [PUTATIVE]	Kinase
9768	1255	CDP-DIACYLGLYCEROLGLYCEROL-3-PHOSPHATE 3-	Transferases
		PHOSPHATIDYLTRANSFERASE [PUTATIVE]	
9775	1256	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9779	1257	PHOSPHOLIPASE [PUTATIVE]	Lipase
9780	1258	PHOSPHOLIPASE [PUTATIVE]	Lipase
9781	1259	PHOSPHOLIPASE [PUTATIVE]	Lipase
9801	1260	BETA-1,3-ENDOGLUCANASE [PUTATIVE]	Glycosylase
9803	1261	PROTEIN KINASE [PUTATIVE]	Kinase, Protein
9815	1262	RIBONUCLEASE, RNS2 IDENTICAL TO	Nuclease
		SP:P42814:RNS2_ARATH; CONTAINS A RIBONUCLEASE	
		T2 FAMILY HISTIDINE ACTIVE SITE SIGNATURE	
		(PDOC00459)[PUTATIVE]	
9822	1263	SERINE PROTEASE [PUTATIVE]	Protease

9825	1264	PROLINE TRANSPORTER 1	Transporter
9829	1265	ISOAMYLASE [PUTATIVE]	Glycosylase
9834	1266	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE	Transferases
		[PUTATIVE]	
9859	1267	ANTHRANILATE N-	Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE	
		[PUTATIVE]	
9861	1268	LIPASE/HYDROLASE GDSL-motif SIMILAR TO APG	Lipase
		PROTEINS; PFAM DOMAIN PF00657[PUTATIVE]	
9863	1269	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9867		POLYGALACTURONASE [PUTATIVE]	Glycosylase
9890	1271	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9894		POTASSIUM TRANSPORTER [PUTATIVE]	Transporter
9896	1273	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9897	1274	INITIATOR TRNA PHOSPHORIBOSYL-TRANSFERASE	Transferases
2000		[PUTATIVE]	
9898	1275	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
0000	1056	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9908	1276	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
0000	. 1077	[PUTATIVE]	Dala
9909	12//	GLYCEROL-3-PHOSPHATE DEHYDROGENASE	Dehydrogenases
9927	1278	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
9963	1279	GLUTAMATE SYNTHASE FERRODOXIN-DEPENDENT ,	Synthase
		3' PARTIAL [PUTATIVE]	
9973	1280	DNA-DIRECTED RNA POLYMERASE 23KD SUBUNIT	Polymerase
		[PUTATIVE]	
9987	1281	PEROXIDASE [PUTATIVE]	Oxidase
9990	1282	CYTOKININ OXIDASE [PUTATIVE]	Oxidase
9993	1283	GLYCEROL-3-PHOSPHATE DEHYDROGENASE	Dehydrogenases

10007	1204	THIOREDOXIN REDUCTASE THE LAST 2 EXONS	172 1
10007	1204		1
		ENCODE THIOREDOXIN. THERE IS AN EST MATCH TO	1
		EXONS 5-7, AND THE DISTANCE BETWEEN EXON 7	1
		AND EXON 8 IS ONLY 90BP. IT IS UNLIKELY THIS IS	
		TWO SEPARATE GENES, BUT MORE LIKELY A HYBRID	
		PROTEIN.[PUTATIVE]	
10021	1285	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10024	1286	POLYGALACTURONASE [PUTATIVE]	Glycosylase
10028	1287	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10030	1288	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10031	1289	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10032	1290	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10040	1291	PHOSPHOLIPASE D	Lipase
10066	1292	PROTEIN KINASE, 5' PARTIAL [PUTATIVE]	Kinase, Protein
10085	1293	COPPER AMINE OXIDASE [PUTATIVE]	Oxidase
10096	1294	PHOSPHOENOLPYRUVATE CARBOXYLASE	Carboxylase
10105	1295	LIPASE [PUTATIVE]	Lipase
10115	1296	CITRATE SYNTHASE [PUTATIVE]	Synthase
10127	1297	RIBOSE PHOSPHATE PYROPHOSPHOKINASE	Kinase
		[PUTATIVE]	
10133	1298	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10141	1299	PECTINESTERASE [PUTATIVE]	Esterase
10145	1300	ISOPROPYLMALATE DEHYDRATASE [PUTATIVE]	Dehydratase
10146	1301	ISOPROPYLMALATE DEHYDRATASE	Dehydratase
10151	1302	3-ISOPROPYLMALATE DEHYDRATASE, SMALL	Dehydratase
		SUBUNIT	
10152	1303	3-ISOPROPYLMALATE DEHYDRATASE, SMALL	Dehydratase
		SUBUNIT	
10160	1304	CARBOXYPHOSPHONOENOLPYRUVATE MUTASE	Mutase
		[PUTATIVE]	j
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10165	1305	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinace Protein
10105	1505	DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Killase, Trotelli
10177	1306	BIOTIN SYNTHASE (BIO B)	Synthase
10189		PEROXIDASE [PUTATIVE]	Oxidase
10197		PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP TYPE	L
10197	1508	[PUTATIVE]	isomerase
10198	1309	ENDOCHITINASE [PUTATIVE]	Chitinase
10199	1310	ENDOCHITINASE [PUTATIVE]	Chitinase
10200	1311	ENDOCHITINASE [PUTATIVE]	Chitinase
10201	1312	ENDOCHITINASE [PUTATIVE]	Chitinase
10202	1313	ENDOCHITINASE [PUTATIVE]	Chitinase
10203	1314	ENDOCHITINASE [PUTATIVE]	Chitinase
10207	1315	BETA-1,3-GLUCANASE, C TERMINAL FRAGMENT C	Glycosylase
ŀ		TERMINAL HOMOLOGY ONLY[PUTATIVE]	
10208	1316	BETA-1,3-GLUCANASE, C TERMINAL FRAGMENT C	Glycosylase
		TERMINAL HOMOLOGY ONLY[PUTATIVE]	
10210	1317	RECEPTOR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10211	1318	RECEPTOR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10212	1319	STEAROYL-ACP DESATURASE	Desaturases
10223	1320	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
10224	1321	PROTEIN KINASE CONTAINS A PROTEIN KINASE DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Kinase, Protein
10225	1322	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
10226		PROTEIN KINASE CONTAINS A PROTEIN KINASE DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Kinase, Protein
10227			Glycosylase
10227			Glycosylase
10228		` · · · · · · · · · · · · · · · · · · ·	
		· · · · · · · · · · · · · · · · · · ·	Glycosylase
10230		-	Glycosylase
10232	1328	METHYL CHLORIDE TRANSFERASE [PUTATIVE]	Transferases
10234	1329	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10246	1330	6,7-DIMETHYL-8-RIBITYLLUMAZINE SYNTHASE	Synthase
İ		PRECURSOR	

10293	1331	HEME A: FARNESYLTRANSFERASE [PUTATIVE]	Transferases
10320	1332	FLAVONOL SYNTHASE [PUTATIVE]	Synthase
10321	1333	TRIACYLGLYCEROL LIPASE [PUTATIVE]	Lipase
10353	1334	PHOSPHATIDATE CYTIDYLYLTRANSFERASE	Transferases
		[PUTATIVE]	
10360	1335	PECTINESTERASE [PUTATIVE]	Esterase
10368	1336	5-ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE (EPSP)	Synthase
		SYNTHASE IDENTICAL TO GB:X06613:ATEPSPS	
10369	1337	NUCLEOTIDE SUGAR EPIMERASE [PUTATIVE]	Epimerase
10372	1338	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10377	1339	FLAVONOL REDUCTASE [PUTATIVE]	Reductase
10381	1340	DIHYDRODIPICOLINATE SYNTHASE [PUTATIVE]	Synthase
10396	1341	PROTEIN KINASE CONTAINS A PROTEIN KINASE	Kinase, Protein
		DOMAIN PROFILE (PDOC00100)[PUTATIVE]	
10425	1342	BETA-AMYLASE [PUTATIVE]	Glycosylase
10448	1343	3-METHYL-2-OXOBUTANOATE HYDROXY-METHYL-	Transferases
		TRANSFERASE	
10479	1344	CYCLIC NUCLEOTIDE-REGULATED ION CHANNEL	Channel
		PROTEIN [PUTATIVE]	
10480	1345	CYCLIC NUCLEOTIDE-REGULATED ION CHANNEL	Channel
		PROTEIN [PUTATIVE]	
10481		CYCLIC NUCLEOTIDE-REGULATED ION CHANNEL	Channel
		PROTEIN [PUTATIVE]	
10508		BETA-KETOACYL-COA SYNTHASE [PUTATIVE]	Synthase
10521	1348	SER/THR PROTEIN KINASE [PUTATIVE]	Kinase, Protein
10522	1349	INORGANIC PYROPHOSPHATASE 3'	Phosphatase
		PARTIAL[PUTATIVE]	
10529	1350	PECTINESTERASE [PUTATIVE]	Esterase
10539	1351	PECTINESTERASE [PUTATIVE]	Esterase
10540	1352	PECTINESTERASE [PUTATIVE]	Esterase
10541	1353	PECTINESTERASE [PUTATIVE]	Esterase
10547	1354	ALCOHOL DEHYDROGENASE [PUTATIVE]	Dehydrogenases
10548	1355	ALCOHOL DEHYDROGENASE [PUTATIVE]	Dehydrogenases
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10549	1356	ALCOHOL DEHYDROGENASE [PUTATIVE]	Dehydrogenases
10550	1357	ALCOHOL DEHYDROGENASE [PUTATIVE]	Dehydrogenases
10553	1358	GALACTINOL SYNTHASE [PUTATIVE]	Synthase
10563	1359	PECTINESTERASE [PUTATIVE]	Esterase
10567	1360	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE [PUTATIVE]	Isomerase
10578	1361	HISTIDINE KINASE IDENTICAL TO GB:D87545[PUTATIVE]	Kinase, Protein
10590	1362	PECTINESTERASE [PUTATIVE]	Esterase
10594	1363	PHOTOLYASE/BLUE-LIGHT RECEPTOR (PHR2)	Receptor
10598	1364	PHOSPHOLIPASE [PUTATIVE]	Lipase
10602	1365	PECTINESTERASE [PUTATIVE]	Esterase
10608	1366	GLUTATHIONE S-TRANSFERASE (GST6) IDENTICAL TO GB:X95295. BASED ON IDENTICAL CDNA HITS, THE TRANSLATION IS NOW 40 AAS LONGER AT THE N-TERMINAL, AND START OF EXON2 IS ALSO CORRECTED.	Transferases
10628	1367	SERINE PROTEASE [PUTATIVE]	Protease
10636		PROTEIN KINASE CONTAINS A PROTEIN KINASE DOMAIN PROFILE (PDOC00100)[PUTATIVE]	Kinase, Protein
10644	1369	PROTEIN KINASE	Kinase, Protein
10645		EXONUCLEASE CONTAINS ZINC-FINGER C2H2-TYPE DOMAIN; SIMILAR TO X.LAEVIS XPMC2 PROTEIN (XPMC2 PREVENTS MITOTIC CATASTROPHE IN FISSION YEAST)[PUTATIVE]	Nuclease
10670		GLYCOGEN SYNTHASE SIMILAR TO GLYCOGEN SYNTHASE PRECURSOR (GRANULE-BOUND STARCH SYNTHASE II) GB:Q43093 FROM [PISUM SATIVUM][PUTATIVE]	Synthase
10671		PEROXIDASE VERY SIMILAR TO PEROXIDASE GB:CAA66963 FROM [ARABIDOPSIS THALIANA][PUTATIVE]	Oxidase
10678		ALDOSE 1-EPIMERASE SHOWS SIMILARITY TO ALDOSE EPIMERASES[PUTATIVE]	Epimerase

10679	1374	PECTATE LYASE SIMILAR TO PECTATE LYASE Lyase
		PRECURSOR GB:P40973 [LILIUM
		LONGIFLORUM][PUTATIVE]
10680	1375	VOLTAGE-DEPENDENT ANION-SELECTIVE CHANNEL Channel
10000	1575	PORIN SIMILAR TO OUTER MITOCHONDRIAL
		MEMBRANE PORIN (VOLTAGE-DEPENDENT ANION-
		SELECTIVE CHANNEL PROTEIN) (VDAC) (POM 34)
		GB:P42055 [SOLANUM TUBEROSUM][PUTATIVE]
10682	1376	PROTEIN KINASE SIMILAR TO PROTEIN KINASE APKIA Kinase, Protein
		GB:Q06548 [ARABIDOPSIS THALIANA][PUTATIVE]
10691	1377	VACUOLAR MEMBRANE ATPASE SUBUNIT GATPase
		(AVMA10) IDENTICAL TO VACUOLAR MEMBRANE
		ATPASE SUBUNIT G (AVMA10) GB:AF181688
		[ARABIDOPSIS THALIANA]
10702	1378	CARBONIC ANHYDRASE, CHLOROPLAST PRECURSOR Anhydrase
		IDENTICAL TO CARBONIC ANHYDRASE,
		CHLOROPLAST PRECURSOR GB:P27140 [ARABIDOPSIS
		THALIANA]
10714	1379	BETA-1,4-N-ACETYLGLUCOSAMINYLTRANSFERASE Transferases
		GB:AAD31053 [ARABIDOPSIS
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
10728	1380	AMINO ACID PERMEASE SIMILAR TO AMINO ACID Transporter
		PERMEASE GB:AAB71468 [ARABIDOPSIS
		THALIANA][PUTATIVE]
10734	1381	ADENYLATE KINASE SIMILAR TO ADK ADENYLATE Kinase
		KINASE GB:S50007 [STREPTOMYCES
		COELICOLOR][PUTATIVE]
10736	1382	PROTEIN KINASE C-TERMINAL REGION SIMILAR TO Kinase, Protein
		PROTEIN KINASES: GB:S71277 [ARABIDOPSIS
		THALIANA], GB:CAB43834 [ARABIDOPSIS THALIANA];
		PFAM HMM HIT: EUKARYOTIC PROTEIN KINASE
		DOMAIN[PUTATIVE]
10750	1383	DEHYDROGENASE CONTAINS PFAM PROFILES: Dehydrogenases
		PF00106 SHORT CHAIN DEHYDROGENASE, PF00678
		SHORT CHAIN DEHYDROGENASE/REDUCTASE C-
		TERMINUS[PUTATIVE]
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10751	1384	URIDYLYL TRANSFERASE-LIKE PROTEINS Transferases
10/51	1501	GB:AAD20075, GB:AAC00631 [ARABIDOPSIS]
		THALIANA]; UNKNOWN PROTEIN SIMILAR TO
10754	1385	ASPARTATE KINASE SIMILAR TO ASPARTATE KINASE Kinase
10754	1363	GB:CAA67376 [ARABIDOPSIS THALIANA][PUTATIVE]
10757	1206	
10/3/	1380	POTASSIUM TRANSPORTER SIMILAR TO POTASSIUM Transporter
		TRANSPORTER GB:AAB87687 [ARABIDOPSIS]
10762	1207	THALIANA][PUTATIVE]  UDP-GLUCOSYL TRANSFERASE SIMILAR TO UDP-Transferases
10762	138/	1
		GLUCOSE GLUCOSYLTRANSFERASE GB:BAA34687, INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE
		GB:Q41819 [ZEA MAYS]; CONTAINS PFAM PROFILE:
		UDP-GLUCORONOSYL AND UDP-GLUCOSYL
		TRANSFERASES[PUTATIVE]
10765	1200	PROTEIN KINASE CONTAINS PFAM PROFILE: Kinase, Protein
10703	1300	EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
10772	1389	PHYTOCHELATIN SYNTHETASE GB:CAA07251 Synthase
10772	1507	[ARABIDOPSIS THALIANA], PFAM HMM HIT:
		TNFR/NGFR CYSTEINE-RICH REGION, UNKNOWN
		PROTEIN SIMILAR TO PUTATIVE
10792	1390	COESTERASE DOMAIN Esterase
10812		STEAROYL-ACYL CARRIER PROTEIN DESATURASE Desaturases
		SIMILAR TO STEAROYL-ACYL CARRIER PROTEIN
		DESATURASE GB:CAA07349 FROM [LINUM
		USITATISSIMUM][PUTATIVE]
10813		STEAROYL-ACYL CARRIER PROTEIN DESATURASE Desaturases
		SIMILAR TO STEAROYL-ACYL CARRIER PROTEIN
		DESATURASE GB:CAA07349 FROM [LINUM
		USITATISSIMUM][PUTATIVE]
10814	1393	STEAROYL-ACYL CARRIER PROTEIN DESATURASE Desaturases
		SIMILAR TO STEAROYL-ACYL CARRIER PROTEIN
		DESATURASE GB:CAA07349 FROM [LINUM
		USITATISSIMUM][PUTATIVE]
10825	1394	ASPARTYL PROTEASE CONTAINS PFAM PROFILE: Protease
		PF00026 EUKARYOTIC ASPARTYL
		PROTEASE[PUTATIVE]

10826	1395	PROTEIN PHOSPHATASE-2C (PP2C) SIMILAR TO Phosphatase
		PUTATIVE PROTEIN PHOSPHATASE-2C (PP2C)
		GB:AAC36699 [MESEMBRYANTHEMUM]
		CRYSTALLINUM][PUTATIVE]
10828	1396	S-ADENOSYLMETHIONINE:2- Transferases
		DEMETHYLMENAQUINONE METHYLTRANSFERASE
		SIMILAR TO S-ADENOSYLMETHIONINE:2-
		DEMETHYLMENAQUINONE METHYLTRANSFERASE
		GB:P32165 [ESCHERICHIA COLI][PUTATIVE]
10832	1397	PROTEIN KINASE CONTAINS PFAM PROFILE: PF00069 Kinase, Protein
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
10839	1398	IAA-AMINO ACID HYDROLASE (ILR1) IDENTICAL TO Hydrolase
		IAA-AMINO ACID HYDROLASE (ILR1) GB:U23794
		[ARABIDOPSIS THALIANA]
10840	1399	PROTEIN KINASE CONTAINS PFAM PROFILES: PF00069 Kinase, Protein
		EUKARYOTIC PROTEIN KINASE DOMAIN, PF00560
		LEUCINE RICH REPEAT (5 COPIES)[PUTATIVE]
10850	1400	N-ACETLYTRANSFERASE CONTAINS PFAM PROFILE: Transferases
		PF00583 ACETYLTRANSFERASE (GNAT)
10057	1401	FAMILY[PUTATIVE]
10857	1401	CELLULOSE SYNTHASE CATALYTIC SUBUNIT Synthase
	ł	SIMILAR TO CELLULOSE SYNTHASE CATALYTIC SUBUNIT GB:AAD40885 FROM [ARABIDOPSIS]
		THALIANA][PUTATIVE]
10871	1402	GLUTATHIONE S-TRANSFERASE IDENTICAL TO Transferases
10071	1402	GLUTATHIONE S-TRANSFERASE GB:AAB09584 FROM
		[ARABIDOPSIS THALIANA]
10906	1403	PHOSPHOLIPASE SIMILAR TO UNKNOWN PROTEIN Lipase
		GB:AAC32238 [ARABIDOPSIS THALIANA], POTENTIAL
		PHOSPHOLIPASE C- SIMILAR TO MULTIPLE
		PHOSPHOLIPASE PROTEINS FROM MYCOBACTERIUM
		SPECIES: GB:CAB06146, GB:CAB06147, GB:AAC18944,
		GB:CAB44656[PUTATIVE]
10919	1404	PEROXIDASE SIMILAR TO PEROXIDASE GB:CAA66966 Oxidase
		[ARABIDOPSIS THALIANA][PUTATIVE]
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10920	1405	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE Transferases
		SIMILAR TO PHOSPHORIBOSYLANTHRANILATE
		TRANSFERASE GB:CAA16616 [ARABIDOPSIS
		THALIANA], PHOSPHORIBOSYLANTHRANILATE
		TRANSFERASE GB:BAA13032 [PISUM
		SATIVUM][PUTATIVE]
10929	1406	SER/THR PROTEIN KINASE DOMAIN, SIMILAR TO Kinase, Protein
		SERINE/THREONINE PROTEIN KINASE PTO
		GB:AAB47421 [LYCOPERSICON ESCULENTUM]
10930	1407	METHIONINE SYNTHASE SIMILAR TO COBALAMIN-Synthase
		INDEPENDENT METHIONINE SYNTHASE GB:AAC50037
] ]		[ARABIDOPSIS THALIANA][PUTATIVE]
10950	1408	SHORT-CHAIN TYPE DEHYDROGENASE/REDUCTASE Dehydrogenase
		SIMILAR TO SHORT-CHAIN TYPE
		DEHYDROGENASE/REDUCTASE GB:Q08632 [PICEA
		ABIES][PUTATIVE]
10952	1409	SHORT-CHAIN TYPE DEHYDROGENASE/REDUCTASE Dehydrogenase
		SIMILAR TO SHORT-CHAIN TYPE
		DEHYDROGENASE/REDUCTASE GB:Q08632 [PICEA
		ABIES][PUTATIVE]
10953	1410	BETA-1,3-GLUCANASE SIMILAR TO BETA-1,3- Glycosylase
		GLUCANASE GB:S12402 [NICOTIANA SP], GB:CAA03908
		[CITRUS SINENSIS], GB:S44364 [LYCOPERSICON
		ESCULENTUM][PUTATIVE]
10963	1411	GLUTAMATE RECEPTOR (GLR1) IDENTICAL TO Receptor
		PUTATIVE GLUTAMATE RECEPTOR (GLR1)
		GB:AF079998 [ARABIDOPSIS THALIANA][PUTATIVE]
10981	1412	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
		LIPASE/ACYLHYDROLASE
11010		ETHYLENE RECEPTOR (EIN4) SIMILAR TO ETHYLENE Receptor
		RECEPTOR GB:AAC31123 [MALUS DOMESTICA],
		IDENTICAL TO PUTATIVE ETHYLENE RECEPTOR
	i	GB:AAD02485 [ARABIDOPSIS THALIANA]; PFAM HMM
	ļ	HIT: RESPONSE REGULATOR RECEIVER DOMAIN,
		SIGNAL C TERMINAL DOMAIN[PUTATIVE]

WO 02/10210 PCT/EP01/09892

- 115 -

11022 1414 PROTEIN KINASE SIMILAR TO HYPOTHETICAL Kinase, Prote PROTEIN WHICH CONTAINS EUKARYOTIC PROTEIN KINASE DOMAIN PF 00069 GB:CAB51834 [ORYZA SATIVA]; PFAM HMM HIT: EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]  11032 1415 RIBOSE 5-PHOSPHATE ISOMERASE IDENTICAL TO Isomerase PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS THALIANA][PUTATIVE]	NS EUKARYOTIC PROTEIN  59 GB:CAB51834 [ORYZA  F: EUKARYOTIC PROTEIN  E]  DMERASE IDENTICAL TO Isomerase  PHOSPHATE ISOMERASE  S THALIANA]; SIMILAR TO
KINASE DOMAIN PF 00069 GB:CAB51834 [ORYZA SATIVA]; PFAM HMM HIT: EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]  11032 1415 RIBOSE 5-PHOSPHATE ISOMERASE IDENTICAL TO Isomerase PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR OB:AAA91161 [ARABIDOPSIS]	G9 GB:CAB51834 [ORYZA G1: EUKARYOTIC PROTEIN G2] DMERASE IDENTICAL TO Isomerase PHOSPHATE ISOMERASE S THALIANA]; SIMILAR TO
SATIVA]; PFAM HMM HIT: EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]  11032 1415 RIBOSE 5-PHOSPHATE ISOMERASE IDENTICAL TO Isomerase PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	F: EUKARYOTIC PROTEIN  E]  DMERASE IDENTICAL TO Isomerase PHOSPHATE ISOMERASE S THALIANA]; SIMILAR TO
KINASE DOMAIN[PUTATIVE]  11032 1415 RIBOSE 5-PHOSPHATE ISOMERASE IDENTICAL TO Isomerase  PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE  GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO  RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413  [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS  ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases  IDENTICAL TO ZETA-CAROTENE DESATURASE  PRECURSOR GB:AAA91161 [ARABIDOPSIS	DMERASE IDENTICAL TO Isomerase PHOSPHATE ISOMERASE S THALIANA]; SIMILAR TO
11032 1415 RIBOSE 5-PHOSPHATE ISOMERASE IDENTICAL TO Isomerase  PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	OMERASE IDENTICAL TO Isomerase PHOSPHATE ISOMERASE S THALIANA]; SIMILAR TO
PUTATIVE RIBOSE 5-PHOSPHATE ISOMERASE GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	PHOSPHATE ISOMERASE S THALIANA]; SIMILAR TO
GB:AAD57010 [ARABIDOPSIS THALIANA]; SIMILAR TO RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	S THALIANA]; SIMILAR TO
RIBOSE 5-PHOSPHATE ISOMERASES: GB:BAA10413 [SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	<u>-</u>
[SYNECHOCYSTIS SP], GB:CAB49687 [PYROCOCCUS ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	
ABYSSI], GB:S22097 [ESCHERICHIA COLI][PUTATIVE]  11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS]	:CAB49687 [PYROCOCCUS
11040 1416 ZETA-CAROTENE DESATURASE PRECURSOR NEARLY Desaturases IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	<u> </u>
IDENTICAL TO ZETA-CAROTENE DESATURASE PRECURSOR GB:AAA91161 [ARABIDOPSIS	
PRECURSOR GB:AAA91161 [ARABIDOPSIS	<b>,</b>
l	91161 [ARABIDOPSIS
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-
11067 1417 SERINE/THREONINE PROTEIN KINASE SIMILAR TO Kinase, Protei	EIN KINASE SIMILAR TO Kinase, Protein
MANY OTHER PUTATIVE PROTEIN	PUTATIVE PROTEIN
KINASES[PUTATIVE]	
11072 1418 BRANCHED-CHAIN AMINO ACID AMINOTRANSFERASE Transferases	ACID AMINOTRANSFERASE Transferases
[PUTATIVE]	
11076 1419 SIGNAL PEPTIDASE MICROSOMAL GB:P28687 FROM Protease	OSOMAL GB:P28687 FROM Protease
[GALLUS GALLUS]	
11079 1420 GLUCOSE AND RIBITOL DEHYDROGENASE HOMOLOG Dehydrogenas	HYDROGENASE HOMOLOG Dehydrogenases
SIMILAR TO GB:AAC60580 FROM [HORDEUM	580 FROM [HORDEUM
VULGARE] SHOWING HOMOLOGIES TO BACTERIAL	OLOGIES TO BACTERIAL
GLUCOSE AND RIBITOL	ID RIBITOL
DEHYDROGENASES[PUTATIVE]	VE]
11114 1421 PECTINESTERASE SIMILAR TO PECTINESTERASE Esterase	R TO PECTINESTERASE Esterase
PRECURSOR GB:Q42920 FROM [MEDICAGO	FROM [MEDICAGO
SATIVA][PUTATIVE]	
11115 1422 PECTINESTERASE SIMILAR TO PECTINESTERASE Esterase	TO PECTINESTERASE Esterase
PPE8B PRECURSOR GB:Q43062 FROM [PRUNUS	
PERSICA][PUTATIVE]	
11117 1423 PROTEIN PHOSPHATASE-2C SIMILAR TO PROTEIN Phosphatase	
The spinding of the spinding o	Q43062 FROM [PRUNUS
PHOSPHATASE-2C GB:AAC36699 FROM	Q43062 FROM [PRUNUS C SIMILAR TO PROTEIN Phosphatase

11144	1424	PECTINACETYLESTERASE SIMILAR TO GB:CAA67728	Esterase
		FROM [VIGNA RADIATA][PUTATIVE]	·
11152	1425	RECEPTOR PROTEIN KINASES: GB:CAB43834	Kinase, Protein
		GB:S71277 [ARABIDOPSIS THALIANA][HYPOTHETICAL	
		PROTEIN SIMILAR TO]	}
11158	1426	ALKYL HYDROPEROXIDE REDUCTASE AND THIOL-	Reductase
		SPECIFIC ANTIOXIDANT FAMILY	
11164	1427	DUAL-SPECIFICITY PROTEIN PHOSPHATASE SIMILAR	Phosphatase
		TO DUAL-SPECIFICITY PROTEIN PHOSPHATASE	
		GB:CAA77232 [ARABIDOPSIS THALIANA][PUTATIVE]	
11173	1428	GUANYLATE KINASE SIMILAR TO GUANYLATE	Kinase
		KINASE (GMK) GB:AAD31506 [SALMONELLA	
		TYPHIMURIUM]; CONTAINS PFAM PROFILE: PF00625	
		GUANYLATE KINASE[PUTATIVE]	
11179	1429	GLYCOSYL TRANSFERASE CONTAINS PFAM PROFILE:	Transferases
		PF01501 GLYCOSYL TRANSFERASE FAMILY	
		8[PUTATIVE]	
11180	1430	PROTEIN PHOSPHATASE-2C (PP2C) SIMILAR TO	Phosphatase
		PROTEIN PHOSPHATASE-2C (PP2C) GB:AAC36699	
		[MESEMBRYANTHEMUM CRYSTALLINUM]; CONTAINS	
		PFAM PROFILE: PF00481 PROTEIN PHOSPHATASE	
		2C[PUTATIVE]	
11183	1431	PROLYL 4-HYDROXYLASE ALPHA SUBUNIT	Hydroxylase
		PRECURSOR GB:P54001 [RATTUS NORVEGICUS]	
		[UNKNOWN PROTEIN SIMILAR TO C-TERMINAL	
		PORTION OF]	
11188	1432	DEHYDROQUINASE SHIKIMATE DEHYDROGENASE	Dehydrogenases
		SIMILAR TO DEHYDROQUINASE SHIKIMATE	
		DEHYDROGENASE GB:S46210 [NICOTIANA	
		TABACUM][PUTATIVE]	
11203	1433	PYRUVATE DEHYDROGENASE KINASE, 3' PARTIAL	Dehydrogenase
		SIMILAR TO PYRUVATE DEHYDROGENASE KINASE	Ì
		GB:AAC97601 FROM [ARABIDOPSIS	
,		THALIANA][PUTATIVE]	
11206	1434	BETA-GLUCOSIDASE, PUTATIVE SIMILAR TO BETA-	Glycosylase
		GLUCOSIDASE GB:AAF23823 FROM [ARABIDOPSIS	
	-	THALIANA]	
	L	L	

WO 02/10210 PCT/EP01/09892

- 117 -

11236	1435	POLYGALACTURONASE, UNKNOWN PROTEIN	Glycosylase
		CONTAINS PFAM PROFILE:PF00295	
11242	1436	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
		PECTIN METHYLESTERASE GB:BAA89480 FROM	_
	,	[SALIX GILGIANA]	
11251	1437	ASPARTATE PHOSPHATASE, HYPOTHETICAL PROTEIN	Phosphatase
		CONTAINS PFAM PROFILE: PF00987 RAPA-LIKE	
		BACTERIAL PROTEIN ASPARTATE PHOSPHATASE	
11254	1438	TRNA PSEUDOURIDINE SYNTHASE SIMILAR TO TRNA	Synthase
		PSEUDOURIDINE SYNTHASE A GB:P07649	
		[ESCHERICHIA COLI][PUTATIVE]	
11260	1439	PECTATE LYASE SIMILAR TO PECTATE LYASE 2	Lyase
		GB:AAF19196 [MUSA ACUMINATA][PUTATIVE]	
11261	1440	UDP-GLUCOSE:STEROL GLUCOSYLTRANSFERASE	Transferases
		UDP-GLUCOSE:STEROL GLUCOSYLTRANSFERASE	_ !
		GB:Z83833 [ARABIDOPSIS THALIANA]	
11266	1441	PROTEIN KINASE SIMILAR TO APKIA PROTEIN KINASE	Kinase, Protein
		GB:Q06548 [ARABIDOPSIS THALIANA]; CONTAINS	
		PFAM PROFILE: PF00069 EUKARYOTIC PROTEIN	
		KINASE DOMAIN[PUTATIVE]	
11272	1442	PURPLE ACID PHOSPHATASE SIMILAR TO PURPLE	Phosphatase
		ACID PHOSPHATASE GB:CAA06921 [IPOMOEA	
		BATATAS][PUTATIVE]	
11275	1443	GLUCAN SYNTHASE SIMILAR TO GLUCAN SYNTHASE	Synthase
		GB:AAD11794 [FILOBASIDIELLA NEOFORMANS VAR.	
11006	1111	NEOFORMANS][PUTATIVE]	** 1 1
11286	1444	GTP CYCLOHYDROLASE I SIMILAR TO GTP	Hydrolase
		CYCLOHYDROLASE I GB:P22288 [RATTUS	*
		NORVEGICUS]; CONTAINS PFAM PROFILE: PF01227 GTP	
11201	1445	CYCLOHYDROLASE I	I Ividuala a
11291	1445	BETA-1,3-GLUCANASE PRECURSOR SIMILAR TO BETA-	nyurolase
		1,3-GLUCANASE PRECURSOR GB:AAD10386 [ORYZA SATIVA]; CONTAINS PFAM PROFILE: PF00332	
11202	1115	GLYCOSYL HYDROLASES FAMILY 17[PUTATIVE] GLUCOSYLTRANSFERASE GB:AAD23884	Transferences
11292	1440		1 1411SICIASES
		[ARABIDOPSIS THALIANA], UNKNOWN PROTEIN	
		SIMILAR TO PUTATIVE	

11311	1447	GLUTAMATE RECEPTOR GB:AAD09173 [ARABIDOPSIS THALIANA] AND PUTATIVE LIGAND-GATED IONIC CHANNEL GB:AAC33237 [ARABIDOPSIS THALIANA], PUTATIVE	
11323	1448	PREPHENATE DEHYDRATASE, P-PROTEIN: CHORISMATE MUTASE, SIMILAR TO P-PROTEIN: CHORISMATE MUTASE, PREPHENATE DEHYDRATASE GB:P43900 [HAEMOPHILUS INFLUENZAE][PUTATIVE]	
11327	1449	RIBULOSE-1,5-BISPHOSPHATE  CARBOXYLASE/OXYGENASE SMALL SUBUNIT N- METHYLTRANSFERASE I SIMILAR TO RIBULOSE-1,5- BISPHOSPHATE CARBOXYLASE/OXYGENASE SMALL SUBUNIT N-METHYLTRANSFERASE I GB:AAC29137 [CHLOROPLAST SPINACIA OLERACEA][PUTATIVE]	Transferases
11342	1450	POLYGALACTURONASE (PGA3) IDENTICAL TO POLYGALACTURONASE (PGA3) GB:AJ005584 (MOL. GEN. GENET. 261 (6), 948-952 (1999))	Glycosylase
11343	1451	POLYGALACTURONASE SIMILAR TO POLYGALACTURONASE GB:CAA05892 [ARABIDOPSIS THALIANA][PUTATIVE]	Glycosylase
11344	1452	POLYGALACTURONASE SIMILAR TO POLYGALACTURONASE GB:CAA05892 [ARABIDOPSIS THALIANA][PUTATIVE]	Glycosylase
11345	1453	EXOPOLYGALACTURONASE IDENTICAL TO EXOPOLYGALACTURONASE GB:X72292 (MOL. GEN. GENET. 261 (6), 948-952 (1999))	Glycosylase
11357	1454	POLYGALACTURONASE SIMILAR TO POLYGALACTURONASE GB:AAC26512 [CUCUMIS MELO]; CONTAINS NON-CONSENSUS AA DONOR SPLICE SITE AT EXON 2[PUTATIVE]	Glycosylase
11383	1455	PROTEIN KINASE CONTAINS PFAM PROFILE: PF00069 I EUKARYOTIC PROTEIN KINASE DOMAIN, PF00560 LEUCINE RICH REPEAT (5 COPIES)[PUTATIVE]	Kinase, Protein
11391	1456	PROTEIN KINASE CONTAINS PFAM PROFILE: PF00069 I EUKARYOTIC PROTEIN KINASE DOMAIN (2 COPIES)[PUTATIVE]	Kinase, Protein

11398	1457	D-ALANINE:D-LACTATE LIGASE GB:AAD41882 Ligase
11350	1437	1
	·	[ENTEROCOCCUS FAECIUM][HYPOTHETICAL PROTEIN
11100	1450	SIMILAR TO]
11402	1458	SERINE/THREONINE PROTEIN KINASE SIMILAR TO Kinase, Protein
		SERINE/THREONINE-SPECIFIC KINASE GB:S68589
		[ARABIDOPSIS THALIANA]; PFAM HMM HITS:
		PUTATIVE SERINE/THREONINE PROTEIN KINASE,
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
11405	1459	BETA-GLUCAN SYNTHASE - REVERSIBLY Synthase
		GLYCOSYLATABLE POLYPEPTIDE SIMILAR TO
	i	REVERSIBLY GLYCOSYLATABLE POLYPEPTIDE
		GB:AAB88408 [PISUM SATIVUM] (POSSIBLE
}		COMPONENT OF GOLGI [PUTATIVE]
11413	1460	MITOCHONDRIAL INNER MEMBRANE PROTEASE Protease
		SUBUNIT 2 SIMILAR TO MITOCHONDRIAL INNER
		MEMBRANE PROTEASE SUBUNIT 2 GB:P46972
		[SACCHAROMYCES CEREVISIAE], IDENTICAL TO
		PUTATIVE SIGNAL PEPTIDASE GB:AAD56314
		[ARABIDOPSIS THALIANA]; PFAM HMM HIT: SIGNAL
	,	PEPTIDASES I[PUTATIVE]
11416	1461	PROTEIN KINASE PFAM HMM HIT: EUKARYOTIC Kinase, Protein
		PROTEIN KINASE DOMAIN; SIMILAR TO PUTATIVE
		RECEPTOR SER/THR PROTEIN KINASE GB:AAD56317
		[ARABIDOPSIS THALIANA][PUTATIVE]
11438	1462	PROTEIN KINASE SIMILAR TO HYPOTHETICAL Kinase, Protein
		PROTEINS GB:AAC13615, GB:CAA18746, GB:AAB81672
]		[ARABIDOPSIS THALIANA]; CONTAINS PFAM PROFILE:
		EUKARYOTIC PROTEIN KINASE DOMAIN[PUTATIVE]
11441	1463	GLUTATHIONE TRANSFERASE SIMILAR TO Transferases
		GLUTATHIONE TRANSFERASE GB:CAA71784 [GLYCINE
		MAX][PUTATIVE]
11455		PECTINACETYLESTERASE SIMILAR TO Esterase
		PECTINACETYLESTERASE PRECURSOR GB:CAA67728
		[VIGNA RADIATA][PUTATIVE]
11456	1465	PECTINACETYLESTERASE SIMILAR TO Esterase
11450		PECTINACETYLESTERASE PRECURSOR GB:CAA67728
		[VIGNA RADIATA][PUTATIVE]

11465	1466	NON-LTR REVERSE TRANSCRIPTASE SIMILAR TO Transcriptase
		PUTATIVE NON-LTR REVERSE TRANSCRIPTASE
		INCLUDING GB:AAD20714 AND
		GB:AAD24831[PUTATIVE]
11472	1467	PHYTOENE DESATURASE GB:P28553 FROM [GLYCINE Desaturases
		MAX][PUTATIVE, OXIDOREDUCTASE SIMILAR TO]
11478	1468	ASCORBATE PEROXIDASE STRONG SIMILARITY TO Oxidase
		ASCORBATE PEROXIDASE GB:CAA56340[PUTATIVE]
11497	1469	PROTEIN KINASE SIMILAR TO PROTEIN KINASE Kinase, Protein
		(APK1A) GB:Q06548 [ARABIDOPSIS THALIANA];
		CONTAINS PFAM PROFILE: PF00069 EUKARYOTIC
		PROTEIN KINASE DOMAIN[PUTATIVE]
11507	1470	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE Lipase
		CONTAINS PFAM PROFILE: PF00657
	·	LIPASE/ACYLHYDROLASE
11515	1471	ENDONUCLEASE III HOMOLOGS: GB:AAD35453, Nuclease
		GB:BAA79061, GB:CAB49586[HYPOTHETICAL PROTEIN
		SIMILAR TO]
11517	1472	URIDYLATE KINASE SIMILAR TO URIDYLATE KINASE Kinase
		GB:CAB13524 [BACILLUS SUBTILIS], GB:P74457
		[SYNECHOCYSTIS PCC6803][PUTATIVE]
11520	1473	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-Isomerase
		TYPE, UNKNOWN PROTEIN PFAM HMM HIT:
11537	1474	LYCOPENE BETA CYCLASE IDENTICAL TO LYCOPENE Cyclase
		BETA CYCLASE GB:AAB53337 [ARABIDOPSIS
		THALIANA]
11542	1475	FATTY ACID ELONGASE 3-KETOACYL-COA SYNTHASE Synthase
		1 SIMILAR TO FATTY ACID ELONGASE 3-KETOACYL-
		COA SYNTHASE 1 GB:AAC99312 [ARABIDOPSIS
11555	1455	THALIANA][PUTATIVE]
11576	1476	DIADENOSINE 5',5"'-P1,P4-TETRAPHOSPHATE Hydrolase
		HYDROLASE SIMILAR TO DIADENOSINE 5',5"-P1,P4-
		TETRAPHOSPHATE HYDROLASE GB:AAC49902
		[LUPINUS ANGUSTIFOLIUS], PFAM HMM HIT:
	: 	BACTERIAL MUTT PROTEIN[PUTATIVE]

11577	1477	MANNOSYLTRANSFERASE GB:BAA28328 Transferases
		[ESCHERICHIA COLI], UNKNOWN PROTEIN C-
		TERMINAL PORTION SIMILAR TO
11581	1478	ABC TRANSPORTER ATPASE SIMILAR TO ABC Transporter
		TRANSPORTER ATPASE GB:AAC68280 [CHLAMYDIA
]		TRACHOMATIS][PUTATIVE]
11584	1479	GALACTOKINASE, 5' PARTIAL SIMILAR TO Kinase
		GALACTOKINASE GB:BAA84705 [MUS
		MUSCULUS][PUTATIVE]
11585	1480	PECTINESTERASE CONTAINS SIMILARITY TO Esterase
		PECTINESTERASE GB:AAB57671 [CITRUS
		SINENSIS][PUTATIVE]
11586	1481	PECTINESTERASE CONTAINS SIMILARITY TO Esterase
		PECTINESTERASE GB:AAB57671 [CITRUS
		SINENSIS][PUTATIVE]
11588	1482	ALPHA-L-ARABINOFURANOSIDASE CONTAINS Glycosylase
		SIMILARITY TO ALPHA-L-ARABINOFURANOSIDASE A
		PRECURSOR GB:P42254 [ASPERGILLUS
		NIGER][PUTATIVE]
11598	1483	ALPHA/BETA HYDROLASE CONTAINS PFAM PROFILE: Hydrolase
		PF00561 ALPHA/BETA HYDROLASE FOLD; PREDICTED
11601	1404	BY GENSCAN[PUTATIVE]
11601	1484	ALPHA-HYDROXYNITRILE LYASE SIMILAR TO ALPHA-Lyase HYDROXYNITRILE LYASE GB:CAA11219 [MANIHOT]
		ESCULENTA][PUTATIVE]
11603	1/105	(1-4)-BETA-MANNAN ENDOHYDROLASE SIMILAR TO Hydrolase
11003	1403	(1-4)-BETA-MANNAN ENDOHYDROLASE SIMILAR TOTTYGIOLASE
		[LYCOPERSICON ESCULENTUM][PUTATIVE]
11604	1486	(1-4)-BETA-MANNAN ENDOHYDROLASE SIMILAR TO Hydrolase
	2.00	(1-4)-BETA-MANNAN ENDOHYDROLASE GB:AAB87859
		[LYCOPERSICON ESCULENTUM][PUTATIVE]
11631	1487	LEUCOANTHOCYANIDIN DIOXYGENASE, PUTATIVE Oxygenases
	,	SIMILAR TO LEUCOANTHOCYANIDIN DIOXYGENASE
		SP:P51093 [VITIS VINIFERA (GRAPE)]
LL		

11645	1488	GLUCOSYL TRANSFERASE, PUTATIVE SIMILAR TO	Transferases
		ZEATIN O-XYLOSYLTRANSFERASE SP:P56725	
		[PHASEOLUS VULGARIS (KIDNEY BEAN) (FRENCH	
		BEAN)]	
11651	1489	PROTEIN PHOSPHATASE 2C (PP2C) IDENTICAL TO	Phosphatase
		PROTEIN PHOSPHATASE 2C (PP2C) GB:P49598	
		[ARABIDOPSIS THALIANA]	
11676	1490	DIGALACTOSYLDIACYLGLYCEROL SYNTHASE	Synthase
		IDENTICAL TO DIGALACTOSYLDIACYLGLYCEROL	
		SYNTHASE GB:AAD42378 [ARABIDOPSIS THALIANA]	
11684	1491	DIHYDRONEOPTERIN ALDOLASE SIMILAR TO	Aldolase
	!	DIHYDRONEOPTERIN ALDOLASE GB:P28823	
	. !	[BACILLUS SUBTILIS][PUTATIVE]	
11714	1492	DNA-3-METHLYADENINE GLYCOSYLASE (MAG)	Glycosylase
		IDENTICAL TO DNA-3-METHLYADENINE	
		GLYCOSYLASE (MAG) SP:Q39147 [ARABIDOPSIS	
		THALIANA (MOUSE-EAR CRESS)]	
11743	1493	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-	Isomerase
	i	TYPE, PUTATIVE CONTAINS PFAM PROFILE:PF00254	
		FKBP-TYPE PEPTIDYL-PROLYL CIS-TRANS ISOMERASE	
11759	1494	CHITINASE BASIC, IDENTICAL TO BASIC CHITINASE	Chitinase
		GB:AAA32769 GI:166666 [ARABIDOPSIS THALIANA]	
		(PLANT PHYSIOL. 93, 907-914 (1990))	
11771		PROTEIN PHOSPHATASE 2C, PUTATIVE CONTAINS	Phosphatase
		PFAM PROFILE: PF00481 PROTEIN PHOSPHATASE 2C	
11784		ZINC TRANSPORTER IDENTICAL TO PUTATIVE ZINC	Transporter
		TRANSPORTER GB:AAC24197 FROM [ARABIDOPSIS	
		THALIANA], ( PROC. NATL. ACAD. SCI. U.S.A. 95 (12),	
		7220-7224 (1998))[PUTATIVE]	
11796	1497	OXIDOREDUCTASES[HYPOTHETICAL PROTEIN	Reductase
		SIMILAR TO]	
11799	1498	OXIDOREDUCTASES[HYPOTHETICAL PROTEIN	Reductase
		SIMILAR TO]	
11816	1499	PROTEIN KINASE, 3' PARTIAL LEUCINE-RICH REPEAT	Kinase, Protein
		TRANSMEMBRANE PROTEIN KINASE 1 GB:AAC27894	}
		FROM [ZEA MAYS][PUTATIVE]	

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11818	1500	ABC TRANSPORTER, PUTATIVE SIMILAR TO ATMRP4	Transporter
		(TRANSPORT OF GLUTATHIONE-CONJUGATES INTO	
		THE VACUOLE) GB:CAA05625 [ARABIDOPSIS	
		THALIANA]; CONTAINS PFAM PROFILE: PF00005 ABC	
		TRANSPORTER	
11819	1501	ABC TRANSPORTER, PUTATIVE SIMILAR TO ATMRP4	Transporter
		(TRANSPORT OF GLUTATHIONE-CONJUGATES INTO	
		THE VACUOLE) GB:CAA05625 [ARABIDOPSIS	
		THALIANA]; CONTAINS PFAM PROFILE: PF00005 ABC	
		TRANSPORTER	
11820	1502	ABC TRANSPORTER, PUTATIVE SIMILAR TO ATMRP4	Transporter
		(TRANSPORT OF GLUTATHIONE-CONJUGATES INTO	
		THE VACUOLE) GB:CAA05625 [ARABIDOPSIS	
		THALIANA]; CONTAINS PFAM PROFILE: PF00005 ABC	
		TRANSPORTER	
11821	1503	SERINE ACETYLTRANSFERASE (SAT-1) IDENTICAL TO	Transferases
		SERINE ACETYLTRANSFERASE (SAT-1) GB:U22964	
		[ARABIDOPSIS THALIANA] (PLANT MOL. BIOL. 30 (5),	
		1041-1049 (1996))	
11843	1504	LOW AFFINITY CALCIUM ANTIPORTER CAX2 ALMOST	Transporter
		IDENTICAL TO GB:AAB05914 FROM [ARABIDOPSIS	
·		THALIANA], CONTAINS PFAM PROFILE: PF00002 7	
}		TRANSMEMBRANE RECEPTOR (SECRETIN FAMILY),	
		AND PF01699 SODIUM/CALCIUM EXCHANGER PROTEIN	
11849	1505	BRASSINOSTEROID RECEPTOR KINASE, PUTATIVE	Kinase, Protein
		SIMILAR TO GB:AAC49810 FROM [ARABIDOSPSIS	ļ
		THALIANA], CONTAINS PFAM PROFILES: PF00560	
		LEUCINE RICH REPEAT (23 COPIES)	·
11850	1506	L-ASCORBATE OXIDASE PRECURSOR, PUTATIVE	Oxidase
		SIMILAR TO GB:Q00624 FROM [BRASSICA NAPUS]	
	!	(PLANT J. 2 (3), 331-342 (1992))	j
11851	1507	POLLEN SPECIFIC PROTEIN, PUTATIVE SIMILAR TO	Oxidase
		GB:CAB59910 FROM [ARABIDOPSIS THALIANA],	
	,	CONTAINS PFAM PROFILES: PF00394 MULTICOPPER	
		OXIDASE	

11868	1508	GLUCAN ENDO-1,3-BETA-GLUCOSIDASE PRECURSOR,	Glycosylase
		PUTATIVE SIMILAR TO GB:P52409 FROM [ TRITICUM	
		AESTIVUM]	
11874	1509	TRANSPORTER PROTEINS[HYPOTHETICAL PROTEIN	Transporter
		CONTAINS SIMILARITY TO]	
11887	1510	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE GB:BAA24694 FROM [ARABIDOPSIS	
		THALIANA]	
11899	1511	BETA-FRUCTOFURANOSIDASE, PUTATIVE SIMILAR TO	Glycosylase
		BETA-FRUCTOFURANOSIDASE 1 GB:S37212 FROM	
		[ARABIDOPSIS THALIANA]	
11900	1512		Glycosylase
11001	1.710	GB:S37212 FROM [ARABIDOPSIS THALIANA]	
11901	1513	·	Hydrolase
		DEPENDENT HYDROLASE GB:AAD18619 FROM	
11925	1514	[CHLAMYDOPHILA PNEUMONIAE] POLYGALACTURONASE SIMILAR TO	Chrosovice
11923	1314	POLYGALACTURONASE GB:CAA11160 FROM	Glycosylase
		[ARABIDOPSIS THALIANA][PUTATIVE]	Į
11926	1515	GTP PYROPHOSPHOKINASE SIMILAR TO GTP.	Kinase
		PYROPHOSPHOKINASE GB:087331 FROM	
		[CORYNEBACTERIUM GLUTAMICUM][PUTATIVE]	
11928	1516	PROTEASE, 5'PARTIAL SIMILAR TO SERINE PROTEASE	Protease
		GB:CAA07250 FROM [LYCOPERSICON	
		ESCULENTUM][PUTATIVE]	
11944	1517	LIPASE/HYDROLASE GDSL-LIKE MOTIF; MYROSINASE-	Lipase
		ASSOCIATED PROTEIN, PUTATIVE SIMILAR TO	ĺ
		GB:CAA71238 FROM [BRASSICA NAPUS], CONATAINS	
		PFAM PROFILE:PF00657	
11954	1518		Esterase
		PECTINESTERASE GB:Q43867 FROM [ARABIDOPSIS	
11000	1.510	THALIANA][PUTATIVE]	
11955	1519	PECTIN METHYLESTERASE SIMILAR TO PECTIN	Esterase
		METHYLESTERASE GB:Q42534 FROM [ARABIDOPSIS]	
		THALIANA][PUTATIVE]	

11959	1520	LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN Kinase, Protein KINASE SIMILAR TO LEUCINE-RICH REPEAT TRANSMEMBRANE PROTEIN KINASE 1 GB:AAC27894
		FROM [ZEA MAYS][PUTATIVE]
11963	1521	DIAMINOPIMELATE DECARBOXYLASE SIMILAR TO Decarboxylase
		DIAMINOPIMELATE DECARBOXYLASE GB:CAB62550
		FROM [ARABIDOPSIS THALIANA][PUTATIVE]
11970	1522	9-CIS-EPOXYCAROTENOID DIOXYGENASE, PUTATIVE Oxygenases
		SIMILAR TO 9-CIS-EPOXYCAROTENOID DIOXYGENASE
		GB:AAF26356 [PHASEOLUS VULGARIS]
11975	1523	TERPENE SYNTHASE-RELATED PROTEIN, PUTATIVE Synthase
.		CONTAINS PFAM PROFILE: PF01397 TERPENE SYNTHASE FAMILY
11977	1524	GERANYLGERANYL PYROPHOSPHATE SYNTHETASE, Synthase
	1324	PUTATIVE SIMILAR TO GERANYLGERANYL
		PYROPHOSPHATE SYNTHETASE GB:P34802
		[ARABIDOPSIS THALIANA]
11978	1525	TERPENE SYNTHASE GB:CAA72074 FROM Synthase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO]
11979	1526	GERANYLGERANYL PYROPHOSPHATE SYNTHASE Synthase
		GGPS3 ALMOST IDENTICAL TO GB:S71231 FROM
11980	1527	[ARABIDOPSIS THALIANA][PUTATIVE] TERPENE SYNTHASE GB:CAA72074 FROM Synthase
11980	1327	[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO]
11981	1528	GERANYLGERANYL PYROPHOSPHATE SYNTHASE Synthase
		GGPS3 ALMOST IDENTICAL TO GB:S71231 FROM
		[ARABIDOPSIS THALIANA][PUTATIVE]
11983	1529	GLUCAN SYNTHASES[HYPOTHETICAL PROTEIN Synthase
		CONTAINS SIMILARITY TO]
12004	1530	CALLOSE SYNTHASE CATALYTIC SUBUNIT-LIKE Synthase
		PROTEIN GB:CAB88264 GI:7630056 FROM
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO]

12010	1531	RECEPTOR-LIKE SERINE/THREONINE KINASE, Kinase, Protein
		PUTATIVE SIMILAR TO RECEPTOR-LIKE
		SERINE/THREONINE KINASE GB:AAC50043 GI:2465923
		FROM [ARABIDOPSIS THALIANA]
12015	1532	DNA-LIGASE ZN-FINGER REGION (REGION ACTS AS A Ligase
		DNA NICK SENSOR) (3 COPIES AT N-TERMINUS)
12020	1533	PHOSPHOENOLPYRUVATE CARBOXYLASE (PPC) Carboxylase
		IDENTICAL TO PHOSPHOENOLPYRUVATE
		CARBOXYLASE (PPC) GB:AF071788 [ARABIDOPSIS
		THALIANA]
12022	1534	GALACTOSYLTRANSFERASE, PUTATIVE CONTAINS Transferases
		PFAM PROFILE: PF01762 GALACTOSYLTRANSFERASE;
		SIMILAR TO AVR9 ELICITOR RESPONSE PROTEIN
		GB:CAA06925 [NICOTIANA TABACUM]
12053	1535	PROTEIN PHOSPHATASE TYPE 2C SIMILAR TO Phosphatase
		PROTEIN PHOSPHATASE TYPE 2C GB:AAD17805 FROM
		[LOTUS JAPONICUS][PUTATIVE]
12088	1536	PHOSPHATIDYLINOSITOL 3- AND 4-KINASE Kinase
		HYPOTHETICAL PROTEIN CONTAINS PFAM PROFILE:
		PF00454
12095	1537	LYSOPHOSPHOLIPASE SIMILAR TO Lipase
		LYSOPHOSPHOLIPASE GB:AAD52700 [SCHISTOSOMA]
		JAPONICUM][PUTATIVE]
12099	1538	ACETYL-COA CARBOXYLASE BIOTIN-CONTAINING Carboxylase
		SUBUNIT SIMILAR TO ACETYL-COA CARBOXYLASE
		BIOTIN-CONTAINING SUBUNIT GB:AAC49114
		[ARABIDOPSIS THALIANA]; CONTAINS PFAM PROFILE:
		PF00364 BIOTIN-REQUIRING ENZYMES[PUTATIVE]
12102	1539	POLYGALACTURONASE SIMILAR TO Glycosylase
		POLYGALACTURONASE GB:BAA88472 [CUCUMIS
		SATIVUS][PUTATIVE]
12110	1540	BETA-1,3-GLUCANASE PRECURSOR SIMILAR TO BETA-Glycosylase
		1,3-GLUCANASE PRECURSOR GB:AAD10386 [ORYZA
	10.4.	SATIVA][PUTATIVE]
12119	1541	PROTEIN KINASE, PUTATIVE SIMILAR TO SOMATIC Kinase, Protein
		EMBRYOGENESIS RECEPTOR-LIKE KINASE
	-	GB:AAB61708 FROM [DAUCUS CAROTA]

12124	1542	GLYCOSYL TRANSFERASES GROUP 1, HYPOTHETICAL	Transferases
		PROTEIN CONTAINS PFAM PROFILE:PF00534	
12133	1543	RECEPTOR KINASE SIMILAR TO RECEPTOR KINASE	Kinase, Protein
		GB:S70769 FROM [ARABIDOPSIS	
		THALIANA][PUTATIVE]	
12166	1544	LIPASE/HYDROLASE GDSL-LIKE MOTIF; PUTATIVE	Lipase
		CONTAINS PFAM PROFILE: PF00657	
		LIPASE/ACYLHYDROLASE	
12181	1545	GLUCOSYLTRANSFERASE SIMILAR TO UDP-GLUCOSE	
		GLUCOSYLTRANSFERASE GB:X77459 [MANIHOT	
		ESCULENTA], UDP-GLYCOSE:FLAVONOID	
		GLYCOSYLTRANSFERASE GB:BAA36411 [VIGNA	
		MUNGO][PUTATIVE]	
12183	1546	SERINE PROTEASE SIMILAR TO HUMAN HTRA SERINE	Protease
		PROTEASE GB:AAC97211 [HOMO SAPIENS][PUTATIVE]	
12184	1547	SERINE PROTEASE SIMILAR TO HUMAN HTRA SERINE	Protease
		PROTEASE GB:AAC97211 [HOMO SAPIENS][PUTATIVE]	
12186	1548	PROTEIN PHOSPHATASE 2C DOMAIN	Phosphatase
12211	1549	PROTEIN PHOSPHATASE, PUTATIVE SIMILAR TO	Phosphatase
		PROTEIN PHOSPHATASE-2C GB:AAC36699 FROM	
		[MESEMBRYANTHEMUM CRYSTALLINUM]	
12216	1550	POLYGALACTURONASE, PUTATIVE SIMILAR TO	Glycosylase
}		ENDOPOLYGALACTURONASE GB:225933 FROM	
		[LYCOPERSICON ESCULENTUM]	
12224	1551	ACYL-COA SYNTHETASE, AMP-BINDING PROTEIN,	Synthase
		PUTATIVE CONTAINS PFAM PROFILE: PF00501 AMP-	
		BINDING ENZYME; SIMILAR TO ACYL-COA	
		SYNTHETASE GB:CAB54055 [PSEUDOMONAS PUTIDA]	
12225	1552	CHITINASE BASIC, PUTATIVE SIMILAR TO BASIC	Chitinase
		CHITINASE GB:CAA78843 [LYCOPERSICON	
		ESCULENTUM] (PLANT MOL. BIOL. 22 (6), 1017-1029	
		(1993))	
12239	1553	,	Esterase
		PECTINESTERASE GB:AAB57669 [CITRUS SINENSIS];	
		CONTAINS PFAM PROFILE: PF01095 PECTINESTERASE	

12240	1554	PEROXIDASE, PUTATIVE SIMILAR TO PEROXIDASE	Oxidase
		GB:AAD37376 [GLYCINE MAX]	
12242	1555	PROTEIN PHOSPHATASE-2C, PUTATIVE SIMILAR TO	Phosphatase
		PROTEIN PHOSPHATASE-2C GB:AAC35951	
		[MESEMBRYANTHEMUM CRYSTALLINUM]; CONTAINS	
		PFAM PROFILE: PF00481 PROTEIN PHOSPHATASE 2C	
12253	1556	NON-LTR RETROELEMENT REVERSE	Transcriptase
		TRANSCRIPTASES[HYPOTHETICAL PROTEIN SIMILAR	
		тој	
12260	1557	PROTEIN PHOSPHATASE SIMILAR TO PROTEIN	Phosphatase
		PHOSPHATASE-2C GB:AAC36698 FROM	
		[MESEMBRYANTHEMUM CRYSTALLINUM][PUTATIVE]	
12266			Methylase
12277	1559	SERINE/THREONINE PROTEIN KINASE, PUTATIVE	Kinase, Protein
		CONTAINS PFAM PROFILE: PF00069 EUKARYOTIC	
		PROTEIN KINASE DOMAIN	
12306	1560	ION CHANNEL PROTEIN FROM [ARABIDOPSIS	Channel
		THALIANA][HYPOTHETICAL PROTEIN CONTAINS	
		SIMILARITY TO]	
12307	1561	ION CHANNEL PROTEIN FROM [ARABIDOPSIS	Channel
		THALIANA][HYPOTHETICAL PROTEIN CONTAINS	
		SIMILARITY TO]	D - 1 - 1
12309	1562	GLUTAMATE DECARBOXYLASE (GAD) (ERT DI)	Decarboxylase
		GB:P54767 [LYCOPERSICON ESCULENTUM]; SIMILAR TO N-TERMINAL PORTION OF	
10212	1562	GLUTAMATE DECARBOXYLASE, PUTATIVE SIMILAR	Decarbovylace
12313	1203	TO GLUTAMATE DECARBOXYLASE GB:Q07346	Decarboxylase
		[PETUNIA X HYBRIDA] (J. BIOL. CHEM. 268 (26), 19610-	
		19617 (1993))	
12316	1564		Phosphatase
12310	1304	GB:CAB63938 FROM [ARABIDOPSIS THALIANA]	
12333	1565	AMIDASE, PUTATIVE SIMILAR TO AMIDASE	AMIDASE
		GB:AAA26183 FROM [RHODOCOCCUS SP.]	
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12361	1566	DIACYLGLYCEROL PYROPHOSPHATE PHOSPHATASE, Phosphatase
		PUTATIVE SIMILAR TO GB:NP_010570 FROM
]		[SACCHAROMYCES CEREVISIAE], CONTAINS PFAM
		PROFILE: PF01569 PHOSPHATASE PAP2 SUPERFAMILY
12366	1567	MANDELATE RACEMASE/MUCONATE LACTONIZING Epimerase
		ENZYME FAMILY, UNKNOWN PROTEIN CONTAINS
		PFAM PROFILE: PF01188
12383	1568	PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT Phosphatase
		CALCINEURIN B SUBUNIT GB:P42322 FROM
		[NAEGLERIA GRUBERI] (GENE 154 (1), 39-45 (1995))
12389	1569	ASPARTYL PROTEASE, CHLOROPLAST NUCLEOID Protease
		DNA-BINDING PROTEIN SIMILAR TO GB:BAA22813
		FROM [NICOTIANA TABACUM] (PLANT CELL 9 (9),
		1673-1682 (1997)), CONTAINS PFAM PROFILE: PF00026
		EUKARYOTIC ASPARTYL PROTEASE[PUTATIVE]
12410	1570	URIDYLATE KINASE, PUTATIVE SIMILAR TO UMP-Kinase
}		KINASE GB:CAB38122 FROM [LACTOCOCCUS LACTIS]
12423	1571	PROTEIN KINASE, PUTATIVE SIMILAR TO SOMATIC Kinase, Protein
		EMBRYOGENESIS RECEPTOR-LIKE KINASE
		GB:AAB61708 FROM [DAUCUS CAROTA]
12442	1572	FLAVANONE-3-HYDROXYLASE GB:Q05965 FROM Hydroxylase
] · ]		[MATTHIOLA INCANA], CONTAINS PFAM PROFILE:
}		PF00671 IRON/ASCORBATE OXIDOREDUCTASE
		FAMILY[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
12443	1573	FLAVONOL SYNTHASE (FLS) GB:Q41452 FROM Synthase
		[SOLANUM TUBEROSUM], CONTAINS PFAM PROFILE:
		PF00671 IRON/ASCORBATE OXIDOREDUCTASE
		FAMILY[HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO]
12445	1574	PHOSPHOSERINE AMINOTRANSFERASE GB:P19689 Transferases
	•	FROM [YERSINIA ENTEROCOLITICA][HYPOTHETICAL
10455	1575	PROTEIN CONTAINS SIMILARITY TO]
12459	1575	TRNA ISOPENTENYL TRANSFERASE, PUTATIVE Transferases
		SIMILAR TO TRNA ISOPENTENYL TRANSFERASE
		GB:AAF00582 FROM [ARABIDOPSIS THALIANA]

10491	1576	PLICOCYL TRANSPERACE, BUTTATIVE CHALLER TO PLICE C
12471	1576	FUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO FUCT Transferases
		C3 PROTEIN GB:CAB52254 FROM [VIGNA RADIATA] (J.
		BIOL. CHEM. (1999) 274 (31), 21830-21839)
12488	1577	CINNAMYL ALCOHOL DEHYDROGENASE IDENTICAL Dehydrogenases
		TO GB:P48523 FROM [ARABIDOPSIS THALIANA]
12501	1578	AMINO ACID PERMEASE, PUTATIVE, 5' PARTIAL Transporter
		CONTAINS PFAM PROFILE: PF00324 AMINO ACID
		PERMEASE
12508	1579	BETA-XYLOSIDASE, PUTATIVE SIMILAR TO BETA-Glycosylase
		XYLOSIDASE A GB:BAA28267 FROM [ASPERGILLUS
		ORYZAE]
12516	1580	RECEPTOR-LIKE PROTEIN KINASE, PUTATIVE SIMILAR Kinase, Protein
		TO LEUCINE-RICH RECEPTOR-LIKE PROTEIN KINASE
		GB:AAC36318 FROM [MALUS DOMESTICA]
12564	1581	GERANYL GERANYL PYROPHOSPHATE SYNTHASE, Synthase
		PUTATIVE SIMILAR TO GERANYL GERANYL
		PYROPHOSPHATE SYNTHASE GB:BAA23157
		[ARABIDOPSIS THALIANA]
12567	1582	RECEPTOR KINASE PROTEIN, PUTATIVE SIMILAR TO Kinase, Protein
		RECEPTOR KINASE GB:AAA33715 [PETUNIA
		INTEGRIFOLIA]
12572	1583	MITOCHONDRIAL CARRIER PROTEIN, PUTATIVE Transporter
		CONTAINS PFAM PROFILE: PF00153 MITOCHONDRIAL
		CARRIER PROTEINS
12597	1584	TETRAACYLDISACCHARIDE 4'-KINASE GB:P27300 Kinase
		[ESCHERICHIA COLI][HYPOTHETICAL PROTEIN
		SIMILAR TO]
12599	1585	PURPLE ACID PHOSPHATASE, PUTATIVE SIMILAR TO Phosphatase
		PURPLE ACID PHOSPHATASE GB:CAA06921 [IPOMOEA
		BATATAS]
12602	1586	PROTEIN KINASE, PUTATIVE CONTAINS PFAM Kinase, Protein
].	,	PROFILE: PF00069 EUKARYOTIC PROTEIN KINASE
		DOMAIN
12627	1587	DNA TOPOISOMERASE VI SUBUNIT B TYPE II Isomerase
		GB:O05207 FROM [SULFOLOBUS SHIBATAE],
		UNKNOWN PROTEIN SIMILAR TO
L		

12662	1588	PHOSPHORIBOSYAMIDOIMIDAZOLE-	Synthase
12002	1560		
		SUCCINOCARBOXAMIDE SYNTHASE, PUTATIVE	
	·	SIMILAR TO PHOSPHORIBOSYLAMIDOIMIDAZOLE-	}
		SUCCINOCARBOXAMIDE SYNTHASE GB:P38025 FROM	
		[ARABIDOPSIS THALIANA]	
12676	1589		Transporter
		[ARABIDOPSIS THALIANA], SIMILAR TO MRP-LIKE	
12685	1590	SERINE/THREONINE-SPECIFIC PROTEIN KINASE,	Kinase, Protein
		PUTATIVE SIMILAR TO SERINE/THREONINE-SPECIFIC	
		PROTEIN KINASE GB:T02731 FROM [ARABIDOPSIS	
		THALIANA]	
12693	1591	IRON/ASCORBATE OXIDOREDUCTASE FAMILY	Reductase
12696	1592	PROTEIN KINASE DOMAIN	Kinase, Protein
12698	1593	ADENYL CYCLASE, PUTATIVE, 3' PARTIAL SIMILAR TO	Cyclase
		ADENYL CYCLASE GB:AAB87670 FROM [NICOTIANA	
		TABACUM]	
12699	1594	ADENYL CYCLASE, PUTATIVE, 5' PARTIAL SIMILAR TO	Cyclase
] ]		ADENYL CYCLASE GB:AAB87670 [NICOTIANA	
		TABACUM]	
12703	1595	1-D-DEOXYXYLULOSE 5-PHOSPHATE SYNTHASE,	Synthase
		PUTATIVE SIMILAR TO 1-D-DEOXYXYLULOSE 5-	
		PHOSPHATE SYNTHASE GB:AAD38941	
		[LYCOPERSICON ESCULENTUM]	
12709	1596	UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D-	Transferases
		GLUCOSYLTRANSFERASE, PUTATIVE SIMILAR TO	
		UDP-GLUCOSE:INDOLE-3-ACETATE BETA-D-	
		GLUCOSYLTRANSFERASE GB:AAB58497	
12717	1597		Isomerase ·
	•	TYPE, PUTATIVE CONTAINS PFAM PROFILE: PF00254	
		FKBP-TYPE PEPTIDYL-PROLYL CIS-TRANS	
		ISOMERASES	
12728	1598	UDP-GLUCOSE GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
	ı	GB:Q40284 FROM [MANIHOT ESCULENTA][PUTATIVE]	
12729	1599	UDP-GLUCOSE GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
		GB:Q40284 FROM [MANIHOT ESCULENTA][PUTATIVE]	
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12730	1600	PEROXIDASE ALMOST IDENTICAL TO GB:CAA66965	Oxidase
		AND GB:CAA67360 FROM [ARABIDOPSIS	
		THALIANA][PUTATIVE]	
12731	1601	UDP-GLUCOSE GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
		GB:Q40284 FROM [MANIHOT ESCULENTA][PUTATIVE]	
12732	1602	UDP-GLUCOSE GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
		GB:Q40284 FROM [MANIHOT ESCULENTA][PUTATIVE]	
12733	1603	UDP-GLUCOSE GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
		GB:Q40284 FROM [MANIHOT ESCULENTA][PUTATIVE]	
12749	1604	SALICYLIC ACID CARBOXYL METHYLTRANSFERASE,	Transferases
	·	PUTATIVE SIMILAR TO GB:AAF00108 FROM [CLARKIA	
		BREWERI]	
12792	1605	ALTERNATIVE OXIDASE 1B PRECURSOR IDENTICAL	Oxidase
		TO GB:023913 FROM [ARABIDOPSIS THALIANA]	
12793	1606	ALTERNATIVE OXIDASE 1A PRECURSOR IDENTICAL	Oxidase
		TO GB:Q39219 FROM [ARABIDOPSIS THALIANA]	
12813	1607	ACETYLTRANSFERASE (GNAT) FAMILY;	Transferases
		HYPOTHETICAL PROTEIN PREDICTED BY	
		GENEMARK.HMM, CONTAINS PFAM PROFILE:PF00583	
		ACETYLTRANSF	
12849	1608	İ	Polymerase
		POLYPEPTIDE (SUBUNIT 5) GB:P46279 [GLYCINE	
		MAX][HYPOTHETICAL PROTEIN SIMILAR TO]	
12875	1609	ETHYLENE RECEPTOR, PUTATIVE (ETR2) SIMILAR TO	Receptor
		ETHYLENE RECEPTOR HOMOLOG GB:AAD31396 FROM	
		[LYCOPERSICON ESCULENTUM], CONTAINS PFAM	
		PROFILE: PF01590 GAF DOMAIN	
12897	1610	PROTEIN PHOSPHATASE GB:AAD17805 FROM [LOTUS	Phosphatase
		JAPONICUS]	
12902		GMC OXIDOREDUCTASE DOMAIN	Reductase
12910	1612	CYANATE LYASE (CYN) IDENTICAL TO CYANASE	Lyase
		(CYN) GB:AB004568 [ARABIDOPSIS THALIANA]	
12918		DIENELACTONE HYDROLASE FAMILY	Hydrolase
12921	1614	DIENELACTONE HYDROLASE FAMILY	Hydrolase

12922	1615	DUAL-SPECIFICITY PROTEIN PHOSPHATASE Phosphatase
		IDENTICAL TO DSPTP1 PROTEIN GB:CAA77232 FROM
		[ARABIDOPSIS THALIANA]
12924	1616	IPP TRANSFERASE, HYPOTHETICAL PROTEIN Transferases
		CONTAINS PFAM PROFILE: PF01715
12926	1617	PROTEIN KINASE GB:AAD24596 FROM [ARABIDOPSIS Kinase, Protein
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR
		TO][PUTATIVE]
12934	1618	XYLOGLUCAN ENDOTRANSGLYCOSYLASE, PUTATIVE Glycosylase
		SIMILAR TO XYLOGLUCAN
]		ENDOTRANSGLYCOSYLASE 1 GB:CAA10231 FROM
		[FAGUS SYLVATICA] (PLANT PHYSIOL.(1999) 119, 1148-
		1148)
12936	1619	RECEPTOR-LIKE KINASE, PUTATIVE SIMILAR TO TMK Kinase, Protein
		GB:CAA69028 FROM [ORYZA SATIVA]
12938	1620	BETA-1,3-GLUCANASE, PUTATIVE SIMILAR TO BETA-Glycosylase
		1,3-GLUCANASE GB:BAA89481 FROM [SALIX
		GILGIANA]
12943	1621	NAD DEPENDENT EPIMERASE, PUTATIVE CONTAINS Dehydratase
[		PFAM PROFILE: PF01370 NAD DEPENDENT
12052	1600	EPIMERASE/DEHYDRATASE FAMILY
12953	1622	BETA-AMYLASE, PUTATIVE SIMILAR TO BETA-Glycosylase
12055	1600	AMYLASE GB:CAB58423 [ARABIDOPSIS THALIANA]
12955	1623	DIHYDROXYACID DEHYDRATASE, PUTATIVE SIMILAR Dehydratase TO DIHYDROXYACID DEHYDRATASE GB:CAA60939
	·	[SACCHAROMYCES CEREVISIAE]
12964	1624	HYDROXYETHYLTHIAZOLE KINASE, PUTATIVE Kinase
12904	1024	SIMILAR TO HYDOXYETHYLTHIAZOLE KINASE (THIM)
		GB:BAA76743 [ESCHERICHIA COLI]
12965	1625	MUCIN CORE 2 BETA 6-N-Transferases
12903	1023	ACETYLGLUCOSAMINYLTRANSFERASE GB:AAA83244
	*	[BOS TAURUS][HYPOTHETICAL PROTEIN SIMILAR TO]
12974	1626	PECTINESTERASE, PUTATIVE CONTAINS PFAM Esterase
12714	1020	PROFILE: PF01095 PECTINESTERASE
L		I NOT IDE. I I VIO/O I DOTINDO I DIGNO

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12983	1627	9-CIS-EPOXYCAROTENOID DIOXYGENASE, PUTATIVE	Oxygenases
		SIMILAR TO GB:CAB10168 FROM [LYCOPERSICON	
		ESCULENTUM] (J. EXP. BOT. 47, 2111-2112 (1997))	
12984	1628	PECTATE LYASE, PUTATIVE SIMILAR TO GB:AAF19196	Lyase
		FROM [MUSA ACUMINATA]	
12986	1629	RECEPTOR KINASE, PUTATIVE SIMILAR TO CLVI	Kinase, Protein
		RECEPTOR KINASE GB:AAB58929 FROM [ARABIDOPSIS	
].		THALIANA]	
12988	1630	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE	Transcriptase
		GB:AAC63678 GI:3738337 FROM [ARABIDOPSIS	
		THALIANA], HYPOTHETICAL PROTEIN, 3' PARTIAL	
		SIMILAR TO PUTATIVE	
12996	1631	BETA-1,3-GLUCANASE, PUTATIVE SIMILAR TO BETA-	Glycosylase
		1,3-GLUCANASE GB:BAA89481 FROM [SALIX	
		GILGIANA]	
13003	1632	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE GB:BAA24694 FROM [ARABIDOPSIS	
		THALIANA]	
13005			Hydrolase
13017	1634	· · · · · · · · · · · · · · · · · · ·	Kinase, Protein
] ]		GB:BAA24694 FROM [ARABIDOPSIS THALIANA]	
		(PLANT CELL PHYSIOL. 38 (3), 248-258 (1997))	
13018	1635	PROTEIN KINASE, PUTATIVE SIMILAR TO	Kinase, Protein
·		GB:BAA24694 FROM [ARABIDOPSIS THALIANA]	
		(PLANT CELL PHYSIOL. 38 (3), 248-258 (1997))	
13022	1636	PROCESSING PEPTIDASE, CHLOROPLAST	Protease
]		THYLAKOIDAL PROCESSING PEPTIDASE, PUTATIVE	
1		SIMILAR TO GB:CAA71502 FROM [ARABIDOPSIS	
10000	1.605	THALIANA]	
13029	1637		Kinase, Protein
]	ļ	IDENTICAL TO PUTATIVE KINASE-LIKE PROTEIN	
		TMKL1 PRECURSOR GB:P33543 FROM [ARABIDOPSIS	
		THALIANA], (PLANT MOL. BIOL. 23 (2), 415-421	1
12020		(1993))[PUTATIVE]	T
13030		PECTATE LYASE, PUTATIVE SIMILAR TO PECTATE	Lyase
		LYASE GB:AAB71208 FROM [FRAGARIA ANANASSA],	j
	•	(PLANT MOL. BIOL. 34 (6), 867-877 (1997))	

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13034	1639	NADPH-FERRIHEMOPROTEIN REDUCTASE	Reductase
		GB:AAF02110 FROM [ARABIDOPSIS	5
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR	
		TO][PUTATIVE]	
13043	1640	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE GB:BAA24694 FROM [ARABIDOPSIS	;}
		THALIANA]	
13061	1641	ENDOXYLOGLUCAN TRANSFERASE, PUTATIVE	Transferases
		SIMILAR TO GB:C49539 FROM [ARABIDOPSIS	
	   	THALIANA] (J. BIOL. CHEM. 268 (34), 25364-25368 (1993))	
13067	1642	ACYL-(ACYL CARRIER PROTEIN) THIOESTERASE	Esterase
		IDENTICAL TO ACYL-(ACYL CARRIER PROTEIN)	
		THIOESTERASE GB:Z36912 [ARABIDOPSIS THALIANA]	
		(ARCH. BIOCHEM. BIOPHYS. 316 (1), 612-618 (1995))	
13070	1643	GLYCOSYL TRANSFERASE, PUTATIVE CONTAINS	
		PFAM PROFILE: PF01501 GLYCOSYL TRANSFERASE	
		FAMILY 8	
13084	1644	REVERSE TRANSCRIPTASE GB:S65812 [ARABIDOPSIS	Transcriptase
		THALIANA][HYPOTHETICAL PROTEIN SIMILAR TO]	
13085	1645	NITRATE TRANSPORTER, PUTATIVE SIMILAR TO	Transporter
		NITRATE TRANSPORTER (NTL1) GB:AAC28086	
·	<u></u>	[ARABIDOPSIS THALIANA]	
13088			Transporter
13096	1647	WALL-ASSOCIATED SERINE/THREONINE KINASE,	
1		PUTATIVE SIMILAR TO WALL-ASSOCIATED KINASE 4	
		GB:CAA08793 FROM [ARABIDOPSIS THALIANA]	
13103		SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE,	Kinase, Protein
		PUTATIVE SIMILAR TO SOMATIC EMBRYOGENESIS	
		RECEPTOR-LIKE KINASE GB:AAB61708 FROM [DAUCUS	1
		CAROTA]	
13129	ł	TERPENE SYNTHASE-RELATED PROTEIN CONTAINS	Synthase
		PFAM PROFILE: PF01397 TERPENE SYNTHASE FAMILY	
13130		TERPENE SYNTHASE-RELATED PROTEIN CONTAINS	Synthase
		PFAM PROFILE: PF01397 TERPENE SYNTHASE FAMILY	

13131	1651	LIMONENE CYCLASE, PUTATIVE, 5' PARTIAL SIMILAR	Cyclase
		TO LIMONENE CYCLASE GB:BAA08367 FROM [PERILLA	
		FRUTESCENS]	
13132	1652	l	Synthase
		LIMONENE CYCLASE GB:BAA08367 FROM [PERILLA	*
		FRUTESCENS], CONTAINS PFAM PROFILE: PF01397	!
		TERPENE SYNTHASE FAMILY	
13165	1653	CELLULASE, PUTATIVE CONTAINS PFAM PROFILE	Hydrolase
		PF00150 CELLULASE (GLYCOSYL HYDROLASE FAMILY	1
		5)	
13166	1654	CELLULASE, PUTATIVE CONTAINS PFAM PROFILE:	Hydrolase
		  PF00150 CELLULASE (GLYCOSYL HYDROLASE FAMILY	
		5)	
13190	1655	ALPHA GALACTOSIDASE GB:AAA73963 [GLYCINE	Glycosylase
		MAX}, UNKNOWN PROTEIN SIMILAR TO	
13214	1656	POLYGALACTURONASE, PUTATIVE SIMILAR TO	Glycosylase
		POLYGALACTURONASE (PG1) GB:AAD46483 [GLYCINE	
		MAX]	
13226	1657	PROTEIN KINASE, PUTATIVE CONTAINS PFAM	Kinase, Protein
		PROFILE: PF00069 EUKARYOTIC PROTĘIN KINASE	
		DOMAIN	
13228	1658	ALPHA-MANNOSIDASE, PUTATIVE SIMILAR TO	Glycosylase
	!	LYSOSOMAL ALPHA-MANNOSIDASE GB:AAC34130	
	1	[HOMO SAPIENS] (HUM. MOL. GENET. 6 (5), 717-726	
		(1997))	
13234	1659	SHORT CHAIN ALCOHOL DEHYDROGENASE	1 ' '
		CONTAINS PFAM PROFILE: PF00106 SHORT CHAIN	
		DEHYDROGENASE; SIMILAR TO SEX DETERMINATION	
		PROTEIN TASSELSEED 2 GB:P50160 [ZEA	
		MAYS][PUTATIVE]	
13235	1660	SHORT CHAIN ALCOHOL DEHYDROGENASE	, ,
		CONTAINS PFAM PROFILE: PF00106 SHORT CHAIN	
		DEHYDROGENASE; SIMILAR TO SEX DETERMINATION	
		PROTEIN TASSELSEED 2 GB:P50160 [ZEA	
153:5		MAYS][PUTATIVE]	
13240	1661	ALPHA/BETA HYDROLASE FOLD	Hydrolase

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13248	1662	SHIKIMATE KINASE SIMILAR TO SHIKIMATE KINASE	
		PRECURSOR GB:CAA45121 [LYCOPERSICON	
		ESCULENTUM][PUTATIVE]	
13254	1663	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN	Kinase, Protein
		KINASE APK1A GB:Q06548 FROM [ARABIDOPSIS	
		THALIANA]	
13271	1664	PEPTIDASE FAMILY M48	Protease
13274	1665	PROTEIN PHOSPHATASE 2C, PUTATIVE SIMILAR TO	Phosphatase
		PROTEIN PHOSPHATASE 2C GB:T09640 FROM	
		[MEDICAGO SATIVA]	
13292	1666	ESTERASE, PUTATIVE SIMILAR TO ESTERASE HDE	Esterase
		GB:BAA82510 FROM [PETROLEUM-DEGRADING	
		BACTERIUM HD-1]	
13302	1667	PECTATE LYASE, PUTATIVE SIMILAR TO PECTATE	Lyase
		LYASE GB:CAA70735 [ZINNIA ELEGANS] (PLANT J. 13	
		(1), 17-28 (1998)); CONTAINS PFAM PROFILE: PF00544	
		PECTATE LYASE	
13326	1668	ALTERNATIVE OXIDASE 1C PRECURSOR IDENTICAL	Oxidase
		TO ALTERNATIVE OXIDASE 1C PRECURSOR GB:O22048	
		FROM [ARABIDOPSIS THALIANA]	
13355	1669	NADPH:QUINONE OXIDOREDUCTASE (NQR) NEARLY	Reductase
		IDENTICAL TO NADPH:QUINONE OXIDOREDUCTASE	
		(NQR) GB:AF145234 [ARABIDOPSIS THALIANA]	
13365	1670	EXONUCLEASE	Nuclease
13366	1671	PECTIN METHYLESTERASE, PUTATIVE SIMILAR TO	Esterase
		PECTIN METHYLESTERASE GB:AAB38794 FROM	}
]		[LYCOPERSICON ESCULENTUM]	
13372	1672	PROTEIN KINASE DOMAIN LRR	Kinase, Protein
12295	1672	NON LTD DEVEDOR TRANSCONDERAGE DURANTE	
13385	1	NON-LTR REVERSE TRANSCRIPTASE, PUTATIVE	ı ranscriptase
	i	SIMILAR TO GB:S65812 FROM [ARABIDOPSIS	
	ì	THALIANA] AND OTHER PUTATIVE NON-LTR REVERSE	
165.55		TRANSCRIPTASES	
13389	ı	PEROXIDASE, PUTATIVE SIMILAR TO PEROXIDASE	Oxidase
		ATP26A GB:CAA72487 GI:1890317 [ARABIDOPSIS	
		THALIANA]	

13394	1675	BETA-1,3 GLUCANASE GB:CAB85903 GI:7414433 [PISUM Glycosylase
		SATIVUM][HYPOTHETICAL PROTEIN SIMILAR TO]
13417	1676	RECEPTOR KINASE, PUTATIVE SIMILAR TO RECEPTOR Kinase, Protein
		KINASE GB:AAD02501 FROM [ARABIDOPSIS
		THALIANA]
13420	1677	PROLYL 4-HYDROXYLASE, PUTATIVE SIMILAR TO Hydroxylase
		PROLYL 4-HYDROXYLASE ALPHA SUBUNIT
		PRECURSOR GB:Q10576 FROM [CAENORHABDITIS
		ELEGANS]
13421	1678	PROLYL 4-HYDROXYLASE, PUTATIVE SIMILAR TO Hydroxylase
ļ		PROLYL 4-HYDROXYLASE ALPHA SUBUNIT
		PRECURSOR GB:Q10576 FROM [CAENORHABDITIS
		ELEGANS]
13424	1679	AAA-TYPE ATPASE GB:AAD31347 GI:4874284 FROM ATPase
		[ARABIDOPSIS THALIANA][HYPOTHETICAL PROTEIN
		SIMILAR TO][PUTATIVE]
13441	1680	PROTEIN KINASE, PUTATIVE SIMILAR TO PROTEIN Kinase, Protein
[		KINASE APK1A GB:Q06548 FROM [ARABIDOPSIS
		THALIANA]
13485	1681	PECTINESTERASE SIMILAR TO PECTINESTERASE Esterase
		PRECURSOR GB:Q43043 [PETUNIA INTEGRIFOLIA];
		CONTAINS PFAM PROFILE: PF01095
13486	1692	PECTINESTERASE[PUTATIVE]
13460	1082	VESICLE TRANSPORT PROTEIN SIMILAR TO V-SNARE Transporter ATVT11A GB:AAF24061 [ARABIDOPSIS]
		ATVTIIA GB:AAF2406I [ARABIDOPSIS] THALIANA][PUTATIVE]
13487	1683	TERPENE SYNTHASE-RELATED PROTEIN CONTAINS Synthase
13.107		PFAM PROFILE: PF01397 TERPENE SYNTHASE FAMILY;
	j	SIMILAR TO EPIDERMAL GERMACRENE C SYNTHASE
		GB:AAC39431 [LYCOPERSICON ESCULENTUM], (+)-
		DELTA-CADINENE SYNTHASE GB:P93665 [GOSSYPIUM]
		HIRSUTUM][PUTATIVE]
13496		TERPENE SYNTHASE-RELATED PROTEIN CONTAINS Synthase
		PFAM PROFILE: PF01397 TERPENE SYNTHASE FAMILY
13497	1685	CHORISMATE MUTASE IDENTICAL TO CHORISMATE Mutase
		MUTASE GB:Z26519 [ARABIDOPSIS THALIANA]

13502	1686	SHORT-CHAIN ALCOHOL DEHYDROGENASE,	Dehydrogenase
		PUTATIVE SIMILAR TO SHORT-CHAIN ALCOHOL	.[
		DEHYDROGENASE GB:AAF04194 [PISUM SATIVUM];	
		CONTAINS PFAM PROFILE: PF00106 SHORT CHAIN	
		DEHYDROGENASE	
13503	1687	SHORT-CHAIN ALCOHOL DEHYDROGENASE,	Dehydrogenase
		PUTATIVE SIMILAR TO SHORT-CHAIN ALCOHOL	
		DEHYDROGENASE GB:AAF04194 [PISUM SATIVUM];	
		CONTAINS PFAM PROFILE: PF00106 SHORT CHAIN	
		DEHYDROGENASE	
13519	1688	TERPENE SYNTHASE, PUTATIVE SIMILAR TO TERPENE	Synthase
		SYNTHASE GB:CAA72074 FROM [ARABIDOPSIS	
		THALIANA], CONTAINS PFAM PROFILE: PF01397	
		TERPENE SYNTHASE FAMILY	
13520	1689	FARNESYLTRANSTRANSFERASE, PUTATIVE	Transferases
		(FRAGMENT) FRAGMENT SIMILAR TO	1
		FARNESYLTRANSTRANSFERASE PRECURSOR	
		GB:T10452 FROM [SINAPIS ALBA]	
13521	1690	GERANYLGERANYL PYROPHOSPHATE SYNTHASE,	Synthase
		PUTATIVE SIMILAR TO GERANYLGERANYL	
		PYROPHOSPHATE SYNTHASE, CHLOROPLAST	
		PRECURSOR GB:P34802 FROM [ARABIDOPSIS	
		THALIANA]	
13532	1691	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE,	Transferases
		PUTATIVE SIMILAR TO ANTHOCYANIN 5-AROMATIC	
		ACYLTRANSFERASE GB:BAA74428 FROM [GENTIANA	
		TRIFLORA]	
13536	1692	FLAVONOL 3-O-GLUCOSYLTRANSFERASE, PUTATIVE	Transferases
		SIMILAR TO FLAVONOL 3-O-GLUCOSYLTRANSFERASE	
		GB:Q43716 FROM [PETUNIA X HYBRIDA]	
13541	1693	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE,	Transferases
	i	PUTATIVE SIMILAR TO ANTHOCYANIN 5-AROMATIC	
		ACYLTRANSFERASE GB:BAA74428 [GENTIANA	
		TRIFLORA]	

13542	1694	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE, Transferases
		PUTATIVE SIMILAR TO ANTHOCYANIN 5-AROMATIC
		ACYLTRANSFERASE GB:BAA74428 [GENTIANA
		TRIFLORA]
13554	1695	ALPHA/BETA HYDROLASE, PUTATIVE CONTAINS Hydrolase
		PFAM PROFILE: PF00561 ALPHA/BETA HYDROLASE
		FOLD
13563	1696	PHYTOCHELATIN SYNTHETASE GB:CAA07251 FROM Synthase
		[ARABIDOPSIS THALIANA] (PLANT PHYSIOL. 117, 1526-
		1526 (1998)), UNKNOWN PROTEIN SIMILAR TO
		PUTATIVEE
13564	1697	NON-LTR RETROELEMENT REVERSE TRANSCRIPTASE Transcriptase
		GB:AAD22368 FROM [ARABIDOPSIS]
		THALIANA][HYPOTHETICAL PROTEIN CONTAINS
		SIMILARITY TO][PUTATIVE]
13576	1698	FRUCTOSE-6-PHOSPHATE 2-KINASE/FRUCTOSE-2,6-Phosphatase
		BISPHOSPHATASE GB:AAF04293, HYPOTHETICAL
		PROTEIN CONTAINS REGIONS SIMILAR TO
13584	1699	DEACETYLVINDOLINE 4-O-ACETYLTRANSFERASE, Transferases
		PUTATIVE SIMILAR TO GB:AAC99311 FROM
		[CATHARANTHUS ROSEUS] (PLANT J. 14 (6), 703-713 (1998))
13595	1700	AMINO ACID TRANSPORTER; UNKNOWN PROTEIN Transporter
13393	1700	CONTAINS PFAM PROFILE: PF01490 TRANSMEMBRANE
		AMINO ACID TRANSPORTER PROTEIN
13612	1701	BETA-MANNAN ENDOHYDROLASE SIMILAR TO (1-4)-Hydrolase
13012	1/01	BETA-MANNAN ENDOHYDROLASE GB:AAB87859
		FROM [LYCOPERSICON ESCULENTUM][PUTATIVE]
13665	1702	VETISPIRADIENE SYNTHASE, PUTATIVE, 5' PARTIAL Synthase
		SIMILAR TO VETISPIRADIENE SYNTHASE
		GB:BAB02386 GI:9294376 FROM [ARABIDOPSIS
		THALIANA]
13685	1703	TERPENE SYNTHASE, PUTATIVE CONTAINS PFAM Synthase
		PROFILE: PF01397 TERPENE SYNTHASE FAMILY

13686	1704	GERANYLGERANYL PYROPHOSPHATE SYNTHASE,	Synthase
		PUTATIVE SIMILAR TO GB:P34802 FROM	
		[ARABIDOPSIS THALIANA] (PLANT PHYSIOL. 104 (4),	
		1469-1470 (1994))	ļ
13695	1705	NON-LTR REVERSE TRANSCRIPTASE, PUTATIVE	Transcriptase
13755	1706	PECTIN METHYLESTERASE [PUTATIVE]	Esterase
13796	1707	PEROXIDASE - LYCOPERSICON ESCULENTUM,	Oxidase
		PIR:S32768[PUTATIVE]	
13824	1708	ARABINOSE KINASE - LIKE PROTEIN PUTATIVE	Kinase
		ARABINOSE KINASE ISA1, ARABIDOPSIS THALIANA,	
		EMBL:ATY14404	
13827	1709	RECEPTOR PROTEIN KINASE -LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE PRKI,	
		LYCOPERSICON ESCULENTUM, PIR:T07865	
13834	1710	]	Glycosylase
		POLYGALACTURONASE, MUSKMELON, PIR:T08213	
13835	1711	ALCOHOL DEHYDROGENASE (ATA1)	Dehydrogenases
13858	1712	SUCROSE SYNTHASE -LIKE PROTEIN SUCROSE	Synthase
		SYNTHASE (SUCROSE-UDP GLUCOSYLTRANSFERASE),	
		ARABIDOPSIS THALINA, SWISSPROT:SUSI_ARATH	
13866	1713	PECTINESTERASE -LIKE PROTEIN PECTINESTERASE	Esterase
ļ		PPE8B PRECURSOR, PRUNUS PERSICA,	
		SWISSPROT:PME_PRUPE	
13868	1714	GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE,	Dehydrogenases
		EQUUS CABALLUS, EMBL:AF097179[PUTATIVE]	
13906	1715	AMINE OXIDASE -LIKE PROTEIN AMINE OXIDASE.	Oxidase
		CANAVALIA LINEATA, EMBL:AF172681	
13918	1716	TRANSPORTER-LIKE PROTEIN MULTIDRUG-EFFLUX	Transporter
		TRANSPORTER BLT, BACILLUS SUBTILIS, PIR:139792	
13919	1717	GLUTATHIONE TRANSFERASE-LIKE PROTEIN	Transferases
		GLUTATHIONE TRANSFERASE, PAPAYA, PIR:T09781	
13921		AMINE OXIDASE, CANAVALIA LINEATA,	Oxidase
		EMBL:AF172681[PUTATIVE]	
13991	l	TRIACYLGLYCEROL LIPASE, PSYCHROBACTER	Lipase
i		IMMOBILIS, PIR:S57275[PUTATIVE]	

14001	1720	TYROSINE-PHOSPHATASE-LIKE PROTEIN PROTEIN-	Phosphatase
		TYROSINE-PHOSPHATASE - SCHIZOSACCHAROMYCES	
		POMBE, PIR:A55446	
14011	1721	PREPHENATE DEHYDRATASE CHLOROPLAST	Dehydratase
		SIMILAR TO BACTERIAL PHEA GENE	·
		PRODUCTS[PUTATIVE]	
14014	1722	HISTONE DEACETYLASE SIMILAR TO MAIZE	
		NUCLEOLAR HISTONE DEACETYLASE (U82815)	
		[PUTATIVE]	
14022	1723	PHOSPHATIDYLCHOLINESTEROL O-	Transferases
		ACYLTRANSFERASE (EC 2.3.1.43) PRECURSOR, MOUSE,	
		PIR:XXMSN[PUTATIVE]	
14023	1724	PROTEINKINASE ATPP -LIKE PROTEIN ATPP PROTEIN,	Kinase, Protein
		BRASSICA NAPUS, EMBL:BNA245479	
14029	1725	MEMBRANE ION ANTIPORTER, AMYCOLATOPSIS	Transporter
		ORIENTALIS, EMBL:AL078635[PUTATIVE]	·
14030	1726	MEMBRANE ION ANTIPORTER, AMYCOLATOPSIS	Transporter
		ORIENTALIS, EMBL:AL078635[PUTATIVE]	
14031	1727	MEMBRANE ION ANTIPORTER, AMYCOLATOPSIS	Transporter
		ORIENTALIS, EMBL:AL078635[PUTATIVE]	
14032	1728	MEMBRANE ION ANTIPORTER, AMYCOLATOPSIS	Transporter
		ORIENTALIS, EMBL:AL078635[PUTATIVE]	
14038	1729	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE	Glycosylase
14045	1730	HIGH-AFFINITY NITRATE TRANSPORTER - LIKE	Transporter
		PROTEIN HIGH-AFFINITY NITRATE TRANSPORTER	
		ACH1, ARABIDOPSIS THALIANA, EMBL:AF019748	
14048	1731	2-PHOSPHOGLYCERATE KINASE - METHANOCOCCUS	Kinase
		JANNASCHII, PIR:A64485[PUTATIVE]	
14072	1732	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE LRK1 - ARABIDOPSIS	
		THALIANA,EMBL:ATLECGENE	
14078	1733	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE LRK1 - ARABIDOPSIS THALIANA,	
		EMBL:ATLECGENE	
·			<del> </del>

14080	1734	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE LRK1 - ARABIDOPSIS THALIANA.
		EMBL:ATLECGENE
14081	1735	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE LRK1 - ARABIDOPSIS THALIANA,
		EMBL:ATLECGENE
14082	1736	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE LRKI - ARABIDOPSIS THALIANA,
,		EMBL:ATLECGENE
14083	1737	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE LRK1 - ARABIDOPSIS THALIANA,
		EMBL:ATLECGENE
14098	1738	TRNA INTRON ENDONUCLEASE - ARABIDOPSIS Nuclease
		THALIANA, EMBL:AB036339[PUTATIVE]
14104	1739	TRANSPORTER PROTEIN[PUTATIVE] Transporter
14105	1740	TRANSPORTER PROTEIN[PUTATIVE] Transporter
14106	1741	PROTEIN KINASE NPK1-RELATED PROTEIN KINASE 2 - Kinase, Protein
		ARABIDOPSIS THALIANA, EMBL:AB000798[PUTATIVE]
14107	1742	TRANSPORTER PROTEIN PEPTIDE TRANSPORT Transporter
		PROTEIN - HORDEUM VULGARE,
		PIR:T04378[PUTATIVE]
14108	1743	TRANSPORTER PROTEIN PEPTIDE TRANSPORT Transporter
		PROTEIN - HORDEUM VULGARE,
		PIR:T04378[PUTATIVE]
14109	1744	TRANSPORTER PROTEIN PEPTIDE TRANSPORT Transporter
		PROTEIN - HORDEUM VULGARE,
14110		PIR:T04378[PUTATIVE]
14110	1745	TRANSPORTER PROTEIN PEPTIDE TRANSPORT Transporter
		PROTEIN - HORDEUM VULGARE,
14111	1716	PIR:T04378[PUTATIVE]
14111	1/46	TRANSPORTER PROTEIN PEPTIDE TRANSPORT Transporter
		PROTEIN - HORDEUM VULGARE,
14110	17.47	PIR:T04378[PUTATIVE]
14118	1747	MAP3K ALPHA 1 PROTEIN KINASE - BRASSICA NAPUS, Kinase, Protein
		EMBL:BNA010091[PUTATIVE]

14125	1748	PROTEIN KINASE - LIKE RECEPTOR-LIKE PROTEIN	Kinase, Protein
		KINASE RLK3, ARABIDOPSIS THALIANA,	
		EMBL:ATH011674	
14131	1749	PROTEIN KINASE - LIKE RECEPTOR-LIKE PROTEIN	Kinase, Protein
		KINASE HOMOLOG, ARABIDOPSIS THALIANA,	
		PIR:T45691	
14149	1750	HISTIDYL-TRNA SYNTHETASE	Synthase
14151	1751	PURPLE ACID PHOSPHATASE PRECURSOR -LIKE	Phosphatase
		PROTEIN PURPLE ACID PHOSPHATASE PRECURSOR	
		PAP1, IPOMOEA BATATAS, EMBL:AF200825	
14153	1752	PROTEIN KINASE - LIKE MEK KINASE ALPHA,	Kinase, Protein
		DICTYOSTELIUM DISCOIDEUM, EMBL:AF093689	
14155	1753	PROTEIN KINASE -LIKE NPK1-RELATED PROTEIN	Kinase, Protein
		KINASE 2 ANP2, ARABIDOPSIS THALIANA,	
		EMBL:AB000798	
14156	1754	DEHYDROGENASE -LIKE PROTEIN ALCOHOL	Dehydrogenases
		DEHYDROGENASE HOMOLOG, RIPENING-RELATED	
		TOMATO, PIR:S39508	j
14163	1755	SERINE/THREONINE-SPECIFIC PROTEIN KINASE (EC	Kinase, Protein
		2.7.1) LRRPK, ARABIDOPSIS THALIANA,	
		PIR:T08975[PUTATIVE]	
14164	1756	PROTEIN (FRAGMENT) SERINE/THREONINE-SPECIFIC	Kinase, Protein
		RECEPTOR PROTEIN KINASE, ARABIDOPSIS	
		THALIANA, PIR:S71277[PUTATIVE]	
14165	1757	PROTEIN (FRAGMENT) SERINE/THREONINE-SPECIFIC	Kinase, Protein
		RECEPTOR PROTEIN KINASE (EC 2.7.1),	
		ARABIDOPSISTHALIANA, PIR:S71277[PUTATIVE]	
14166	1758	SERINE/THREONINE-SPECIFIC RECEPTOR PROTEIN	Kinase, Protein
		KINASE (EC 2.7.1), ARABIDOPSISTHALIANA,	
		PIR:S71277[PUTATIVE]	
14167	1759	SERINE/THREONINE-SPECIFIC PROTEIN KINASE (EC	Kinase, Protein
		2.7.1) LRRPK, ARABIDOPSIS THALIANA,	
		PIR:T08975[PUTATIVE]	
14168		RECEPTOR PROTEIN KINASE -LIKE RECEPTOR-LIKE	Kinase, Protein
		PROTEIN KINASE PRECURSOR, MADAGASCAR	
		PERIWINKLE, PIR:T10060	

14172	1761	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE - CATHARANTHUS
		ROSEUS, EMBL:Z73295
14173	1762	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG LIGHT Kinase, Protein
		REPRESSIBLE RECEPTOR PROTEIN KINASE -
		ARABIDOPSIS THALIANA, PIR:S71277
14174	1763	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG LIGHT Kinase, Protein
		REPRESSIBLE RECEPTOR PROTEIN KINASE -
		ARABIDOPSIS THALIANA, PIR:S71277
14176	1764	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG LIGHT Kinase, Protein
		REPRESSIBLE RECEPTOR PROTEIN KINASE -
	,	ARABIDOPSIS THALIANA, PIR:S71277
14179	1765	LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE - Kinase, Protein
		ARABIDOPSIS THALIANA, PIR:S71277[PUTATIVE]
14188	1766	GIBBERELLIN 20-OXIDASE, SEVERAL OXIDASES, Oxidase
		MAINLY [PUTATIVE]
14189	1767	GIBBERELLIN 20-OXIDASE, SEVERAL OXIDASES, Oxidase
		MAINLY [PUTATIVE]
14196	1768	GLUCOSIDASE-LIKE PROTEIN GLUCAN ENDO-1,3-Glycosylase
		BETA-GLUCOSIDASE PRECURSOR - TRITICUM
		AESTIVUM, SWISSPROT:E13B_WHEAT
14204	1769	GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
		GLUCOSE GLUCOSYLTRANSFERASE - ARABIDOPSIS
		THALIANA, EMBL:AB016819
14205	1770	GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
1		GLUCOSE GLUCOSYLTRANSFERASE - ARABIDOPSIS
		THALIANA, EMBL:AB016819
14206	1771	GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
	ĺ	GLUCOSE GLUCOSYLTRANSFERASE - ARABIDOPSIS
		THALIANA, EMBL:AB016819
14207		GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
	1	GLUCOSE GLUCOSYLTRANSFERASE - ARABIDOPSIS
		THALIANA, EMBL:AB016819

14208	1773	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN Transferases
		GLUCURONOSYL TRANSFERASE HOMOLOG,
		RIPENING-RELATED - LYCOPERSICON
		ESCULENTUM,PIR2:S39507
14209	1774	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN Transferases
		GLUCURONOSYL TRANSFERASE HOMOLOG,
		RIPENING-RELATED - LYCOPERSICON
		ESCULENTUM,PIR2:S39507
14211	1775	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN Transferases
·		GLUCURONOSYL TRANSFERASE HOMOLOG,
		RIPENING-RELATED - LYCOPERSICON
		ESCULENTUM,PIR2:S39507
14213	1776	CHLOROPLAST IMPORT-ASSOCIATED CHANNEL Channel
		PROTEIN HOMOLOG CHLOROPLAST IMPORT-
		ASSOCIATED CHANNEL IAP75 - PISUM
		SATIVUM,PIR2:S55344
14215	1777	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE - ARABIDOPSIS
		THALIANA,PID:E1188577
14223	1778	SUBTILISIN-LIKE PROTEINASE CUCUMISIN (SERINE Protease
		PROTEASE,EC 3.4.21.25) PRECURSOR - CUCUMIS
		MELO,PIR2:A55800
14224	1779	SUBTILISIN-LIKE PROTEINASE CUCUMISIN (SERINE Protease
		PROTEASE,EC 3.4.21.25) PRECURSOR - CUCUMIS
		MELO,PIR2:A55800
14229	1780	COPPER TRANSPORT PROTEIN - LIKE PROTEIN Transporter
		COPPER TRANSPORTER PROTEIN ARABIDOPSIS
		THALIANA,PID:G1082054
14239	1781	BETA-D-GLUCAN EXOHYDROLASE - LIKE PROTEIN Hydrolase
		BETA-D-GLUCAN EXOHYDROLASE, NICOTIANA
		TABACUM, TREMBL:AB017502_1
14240	1782	BETA-D-GLUCAN EXOHYDROLASE - LIKE PROTEIN Hydrolase
		BETA-D-GLUCAN EXOHYDROLASE, NICOTIANA
145 15	1500	TABACUM, TREMBL:AB017502_1
14243	1783	BETA-D-GLUCAN EXOHYDROLASE - LIKE PROTEIN Hydrolase
		BETA-D-GLUCAN EXOHYDROLASE, NICOTIANA
<u></u>		TABACUM, TREMBL:AB017502_1

14244	1784	BETA-D-GLUCAN EXOHYDROLASE - LIKE PROTEIN	Hydrolase
		BETA-D-GLUCAN EXOHYDROLASE, NICOTIANA	
		TABACUM, TREMBL:AB017502_1	
14248	1785	RECEPTOR KINASE-LIKE PROTEIN RECEPTOR KINASE-	Kinase, Protein
		LIKE PROTEIN (XA21), ORYZA LONGISTAMINATA,	
		U72725	
14250	1786	RECEPTOR PROTEIN KINASE - LIKE PROTEIN PROTEIN	Kinase, Protein
		KINASE XA21 RECEPTOR TYPE PRECURSOR, ORYZA	
		SATIVA, PIR:A57676	
14258	1787	1-AMINOCYCLOPROPANE-1-CARBOXYLIC ACID	Oxidase
		OXIDASE - LIKE PROTEIN ACC OXIDASE, BRASSICA	
		OLERACEA, X81628	
14274	1788	11BETA-HYDROXYSTEROID DEHYDROGENASE -	Dehydrogenases
		HOMO SAPIENS,PIR1:DXHUBH[PUTATIVE]	
14275	1789	11BETA-HYDROXYSTEROID DEHYDROGENASE -	Dehydrogenases
		RATTUS NORVEGICUS,PIR1:DXRTBH[PUTATIVE]	
14277	1790	PECTINESTERASE HOMOLOG - PINUS	Esterase
		RADIATA,PIR2:T08112[PUTATIVE]	
14279	1791	PECTINESTERASE-LIKE PROTEIN PECTINESTERASE	Esterase
		(EC 3.1.1.11) - CITRUS SINENSIS,PID:G2098709	
14284	1792	NITRIC OXIDE SYNTHASE, PROTEIN BR-1 PROTEIN -	Synthase
·		HELIX POMATIA,PID:E234055[PUTATIVE]	
14293	1793	ENDOCHITINASE-LIKE PROTEIN BASIC	Chitinase
		ENDOCHITINASE CHB4 PRECURSOR - BRASSICA	
		NAPUS, SWISSPROT:CHI4_BRANA	
14297		RECEPTOR KINASE-LIKE PROTEIN PROTEIN KINASE	Kinase, Protein
		XA21 - ORYZA SATIVA, PIR:A57676	
14306		PECTINESTERASE - ARABIDOPSIS THALIANA,	Esterase .
		PIR:S51370[PUTATIVE]	
14318		ABC-TYPE TRANSPORT PROTEIN-LIKE PROTEIN ATP	Transporter
		BINDING CASSETTE TRANSPORTER ABC2, HOMO	
		SAPIENS ,PIR2:B54774	
14334		PEPTIDE TRANSPORTER PEPTIDE TRANSPORTER	Transporter
		(PTR1), HORDEUM VULGARE, AF023472[PUTATIVE]	

14339	1798	CYCLIC NUCLEOTIDE-GATED CHANNEL CYCLIC	Channel
		NUCLEOTIDE-GATED CATION CHANNEL	,
		ARABIDOPSISTHALIANA, AF067798[PUTATIVE]	
14349	1799	GLYCINE-TRNA LIGASE PRECURSOR, CHLOROPLAST	Ligase
		(EDD1)	
14371	1800	LIPASE - LIKE PROTEIN LIPASE ARAB-1, ARABIDOPSIS	Lipase
		THALIANA, PIR2:S68410	
14378	1801	DEOXYCYTIDYLATE DEAMINASE - HOMO SAPIENS,	Deaminase
		PIR:I55434[PUTATIVE]	
14382	1802	ENDOXYLOGLUCAN TRANSFERASE-LIKE PROTEIN	Transferases
		EXGT1 (ENDOXYLOGLUCAN TRANSFERASE) - PISUM	
		SATIVUM, EMBL:AB015428	
14397	1803	GLUTAMATE-1-SEMIALDEHYDE AMINOTRANSFERASE	Transferases
14407	1804	POLY A POLYMERASE, C-TERMINUS	Polymerase
		POLYADENYLATE-BINDING PROTEINS[PUTATIVE]	
14419	1805	ENDO-POLYGALACTURONASE - LIKE PROTEIN ENDO-	Glycosylase
		POLYGALACTURONASE, ARABIDOPSI THALIANA,	
		GB:CAA05525	
14424	1806	RNA POLYMERASE III SUBUNIT - LIKE PROTEIN RNA	Polymerase
		POLYMERASE III SUBUNIT, HOMO SAPIENS,	
	•	GB:CAB41919	
14435	1807	PEROXIDASE	Oxidase
14436	1808	PEROXIDASE	Oxidase
14440	1809	PYRUVATE KINASE -LIKE PROTEIN VARIOUS	Kinase
		PYRUVATE KINASES FROM PROCARYOTES	
14446	1810	PECTINESTERASE - LIKE PROTEIN PECTINESTERASE,	Esterase
		CITRUS SINENSIS,GB:AAB57670	
14455	1811	SUGAR TRANSPORTER 2, DROSOPHILA	Transporter
		MELANOGASTER, EMBL:AF199484[PUTATIVE]	
14460	1812	6-PHOSPHOGLUCONOLACTONASE - LIKE PROTEIN 6-	LACTONASE
		PHOSPHOGLUCONOLACTONASE (6PGL ), HOMO	
		SAPIENS, EMBL:HSA243972	
14491	1813	RECEPTOR PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		CLAVATA1 RECEPTOR KINASE, ARABIDOPSIS	
		THALIANA, EMBL:ATU96879	
14520	1814	PEROXIDASE ATP21A	Oxidase
L			

14545	1815	FLAVONOL SYNTHASE - LIKE PROTEIN SRGI PROTEIN,	Synthase
		ARABIDOPSIS THALIANA, PIR:S44261	
14547	1816	1	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE (RKL1),	}
		ARABIDOPSIS THALIANA, EMBL:AF084034	
14551	1817		Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE -	
		LIKE PROTEIN ANTHRANILATE N-	
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	
		DIANTHUS CARYOPHYLLUS	
14552	1818	ANTHRANILATE N-	Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE -	
		LIKE PROTEIN ANTHRANILATE N-	
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	
		DIANTHUS CARYOPHYLLUS, PIR:T10717	
14553	1819	ANTHRANILATE N-	Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	
		DIANTHUS CARYOPHYLLUS, PIR:T10711[PUTATIVE]	
14554	1820	ANTHRANILATE N-	Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE -	
		LIKE PROTEIN ANTHRANILATE N-	
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	
		DIANTHUS CARYOPHYLLUS, PIR:T10719	
14563	1821	MYOSIN HEAVY CHAIN KINASE B, DICTYOSTELIUM	Kinase, Protein
		DISCOIDEUM, GB:U90946[PUTATIVE]	
14573	1822	REVERSE TRANSCRIPTASE FRAGMENTS[PUTATIVE]	Transcriptase
14576	1823	PHOSPHOGLYCERATE MUTASES[PUTATIVE]	Mutase
14580			Dehydrogenase
		SAPIENS, EMBL:AF078850[PUTATIVE]	7 0
14598		-	Transferases
		PROTEIN UTP-GLUCOSE GLUCOSYLTRANSFERASE,	
		MANIHOT ESCULENTA, PIR:S41951	
14600		UDP-GLUCOSE:(GLUCOSYL) LPS ALPHA1,3-	Transferases
		GLUCOSYLTRANSFERASE WAAO, E.COLI,	
		EMBL:AF019746[PUTATIVE]	

14622	1000	DEDOVIDACE LIVE PROTERY DEDOVIDACE AFROM	10
14623	1827	PEROXIDASE-LIKE PROTEIN PEROXIDASE ATP6A	-Oxidase
		ARABIODOPSIS THALIANA, EMBL:X98774	
14645	1828	STEROID SULFOTRANSFERASE 2 - BRASSICA NAPUS,	Transferases
		EMBL:AF000306[PUTATIVE]	
14648	1829	FLAVANONE 3-HYDROXYLASE (FH3)	Hydroxylase
14661	1830	PROTEIN PHOSPHATASE 2C -LIKE PROTEIN PROTEIN	Phosphatase
		PHOSPHATASE 2C HOMOLOG, MESEMBRYANTHEMUM	
		CRYSTALLINUM, EMBL:AF097667	
14671	1831	PROTEIN PHOSPHATASE 2C-LIKE PROTEIN PROTEIN	Phosphatase
1 1	•	PHOSPHATASE-2C, MESEMBRYANTHEMUM	
		CRYSTALLINUM, EMBL:AF075580	
14672	1832	GLUTAMATE RECEPTOR PUTATIVE GLUTAMATE	Receptor
		RECEPTOR GLR2, ARABIDOPSIS THALIANA,	
		EMBL:AF079999[PUTATIVE]	
14673	1833	SUGAR TRANSPORTER-LIKE PROTEIN SUGAR	Transporter
		TRANSPORTER, ARABIDOPSIS THALIANA,	
		EMBL:Z50752	
14679	1834	RECEPTOR-PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
. [		RECEPTOR-LIKE PROTEIN KINASE, CATHARANTHUS	
		ROSEUS, PIR:T10060	
14692	1835	SHORT-CHAIN ALCOHOL DEHYDROGENASE-LIKE	Dehydrogenase
		PROTEIN SHORT-CHAIN ALCOHOL DEHYDROGENASE -	
		ZEA MAYS,PIR:A47542	
14695	1836	PROTEIN KINASE PK1, RECEPTOR-LIKE - ZEA MAYS,	Kinase, Protein
		PIR:S33532[PUTATIVE]	
14706	1837	CHLOROPHYLL SYNTHETASE	Synthase
14710	1838	CA2+/H+-EXCHANGING PROTEIN-LIKE ARABIDOPSIS	Transporter
		THALIANA HIGH AFFINITY CALCIUM ANTIPORTER	
		CAX1 ENCODED BY GENBANK ACCESSION NUMBER	
		U57411	j
14717	1839	MYOSIN HEAVY CHAIN KINASE B - DICTYOSTELIUM	Kinase, Protein
		DISCOIDEUM, PID:G1903458[PUTATIVE]	
14721	1840	WAX SYNTHASE-LIKE PROTEIN WAX SYNTHASE -	Synthase
		SIMMONDSIA CHINENSIS, PID:G5020219	
14729	1841	DNA POLYMERASE I -BACILLUS	Polymerase
		STEAROTHERMOPHILUS,PIR2:S70368[PUTATIVE]	
	1		

14740	1842	BETA-KETOACYL-COA SYNTHASE LIKE PROTEIN	Synthase
		  BETA-KETOACYL-COA SYNTHASE - SIMMONDSIA	} `
		CHINENSIS,PID:G1045614	]
14750	1843	PSEUDOURIDINE SYNTHASE, PUTATIVE SIMILAR TO	Synthase
		RIBOSOMAL LARGE SUBUNIT PSEUDOURIDINE	1
		SYNTHASE D SP:P33643 [ESCHERICHIA COLI]	
14777	1844	PROTEIN KINASE-LIKE PROTEIN - ARABIDOPSIS	Kinase, Protein
		THALIANA (F15B8.110),PIR2:T06750[PUTATIVE]	
14784	1845	BETA-FRUCTOFURANOSIDASE	Glycosylase
14786	1846	PHOSPHATE ACTYLTRANSFERASE,	Transferases
		STAPHYLOCOCCUS AUREUS,	
		EMBL:SAU271496[PUTATIVE]	
14796	1847	CARBONIC ANHYDRASE (CAHI)	Anhydrase
14802	1848	PURPLE ACID PHOSPHATASE-LIKE PROTEIN PURPLE	Phosphatase
		ACID PHOSPHATASE PRECURSOR, PHASEOLUS	
		VULGARIS, EMBL:PVPAPHOSP	
14809	1849	VACUOLAR SORTING RECEPTOR HOMOLOG/ATELPI	Receptor
		and SPOT 3 PROTEIN	
14812	1850	MONODEHYDROASCORBATE REDUCTASE (NADH) -	Reductase
		LIKE PROTEIN MONODEHYDROASCORBATE	
		REDUCTASE (NADH), LYCOPERISON ESCULENTUM,	
		PIR:T06407	
14825	1851	PROTEIN KINASE, ARABIDOPSIS	Kinase, Protein
		THALIANA[PUTATIVE]	
14832	1852	BETA-GALACTOSIDASE PRECURSOR. LYCOPERSICON	Glycosylase
		ESCULENTUM, GB:P48980[PUTATIVE]	
14838	1853	CAFFEIC ACID O-METHYLTRANSFERASE - LIKE	Transferases
		PROTEIN CAFFEIC ACID O-METHYLTRANSFERASE	
		(HOMT1), POPULUS KITAKAMIENSIS,	
		EMBL:PKHOMT1A	
14839	1854	•	Transferases
		GLUCOSYLTRANSFERASE IS10A, SALICYLATE-	Ì
		INDUCED, NICOTIANA TABACUM, PIR:T03745	
14840			Transferases
		GLUCOSYLTRANSFERASE IS5A, NICOTIANA	
, ,		TABACUM, PIR:T03747	1

**WO** 02/10210

PROTEIN MTN6, MEDICAGO TRUNCATULA, EMBL:MTY18225  14842 1857 NODULIN / GLUTAMATE-AMMONIA LIGASE - LIKE Ligase PROTEIN MTN6 - NODULIN 6, MEDICAGO TRUNCATULA, EMBL:MET133118  14843 1858 PECTATE LYASE -LIKE PROTEIN PECTATE LYASE, Lyase MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote LECTIN KINASE 3, ARABIDOPSIS THALIANA, GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN EPIMERASE DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459	
14842 1857 NODULIN / GLUTAMATE-AMMONIA LIGASE - LIKE Ligase PROTEIN MTN6 - NODULIN 6, MEDICAGO TRUNCATULA, EMBL:MET133118  14843 1858 PECTATE LYASE -LIKE PROTEIN PECTATE LYASE, Lyase MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote LECTIN KINASE 3, ARABIDOPSIS THALIANA, GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN Epimerase DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	
PROTEIN MTN6 - NODULIN 6, MEDICAGO TRUNCATULA, EMBL:MET133118  14843 1858 PECTATE LYASE -LIKE PROTEIN PECTATE LYASE, Lyase MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote LECTIN KINASE 3, ARABIDOPSIS THALIANA, GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN EPIMERASE DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	1
TRUNCATULA, EMBL:MET133118  14843 1858 PECTATE LYASE -LIKE PROTEIN PECTATE LYASE, Lyase  MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease  CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote  LECTIN KINASE 3, ARABIDOPSIS THALIANA,  GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas  CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN EPIMERASE  DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS  FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	14842
14843 1858 PECTATE LYASE -LIKE PROTEIN PECTATE LYASE, Lyase  MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote LECTIN KINASE 3, ARABIDOPSIS THALIANA, GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN Epimerase DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	
MUSA ACUMINATA, EMBL:MAPEL  14851 1859 CLP ENDOPEPTIDASE ATP-BINDING CHAIN C, Protease CHLAMYDIA PNEUMONIAE, PIR:G72079[PUTATIVE]  14862 1860 RECEPTOR LECTIN KINASE -LIKE PROTEIN RECEPTOR Kinase, Prote LECTIN KINASE 3, ARABIDOPSIS THALIANA, GB:U93161  14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN Epimerase DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	
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14869 1861 LYSINE DECARBOXYLASE (ECORLD), EIKENELLA Decarboxylas CORRODENS, EMBL:U89166[PUTATIVE]  14882 1862 DIAMINOPIMELATE EPIMERASE - LIKE PROTEIN Epimerase DIAMINOPIMELATE EPIMERASE, PSEUDOMONAS FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	
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FLUORESCENS, PIR:T10459  14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	14882
14886 1863 INORGANIC PYROPHOSPHATASE -LIKE PROTEIN Phosphatase	
	14886
INORGANIC PYROPHOSPHATASE, SOLANUM	j
TUBEROSUM, PIR:T07399	14006
14896 1864 NA(+)/H(+) ANTIPORTER[PUTATIVE] Transporter	
14905 1865 SERINE/THREONINE-SPECIFIC KINASE LIKE PROTEIN Kinase, Protei	14905
SERINE/THREONINE-SPECIFIC KINASE (EC 2.7.1)	İ
PRECURSOR - ARABIDOPSIS THALIANA, PIR:S68589  14908 1866 PROTEIN KINASE-LIKE PROTEIN WALL-ASSOCIATED Kinase, Protei	14000
14908 1866 PROTEIN KINASE-LIKE PROTEIN WALL-ASSOCIATED Kinase, Protei KINASE 4 - ARABIDOPSIS THALIANA, EMBL:ATH9695	14908
14914 1867 URACIL PHOSPHORIBOSYLTRANSFERASE-LIKE Transferases	14014
PROTEIN URACIL PHOSPHORIBOSYLTRANSFERASE -	14714
NICOTIANA TABACUM, PIR:T03969	1
	1
Denyulogenas	14915
14919 1869 GLYOXAL OXIDASE (GLX2) - PHANEROCHAETE Oxidase	14915
CHRYSOSPORIUM, EMBL:L47287[PUTATIVE]	
14927 1870 PROTEIN KINASE -LIKE PROTEIN PROTEIN KINASE Kinase, Protei	
APK1, ARABIDOPSIS THALIANA, PIR:S28615	14919

14933	1971	FRUCTOKINASE - LIKE PROTEIN FRUCTOKINASE.	17:
14933	10/1	LYCOPERSICON ESCULENTUM, EMBL:LEU62329	, Kinase
14040	1070	I	
14948	18/2	ESTERASE, PSEUDOMONAS FLUORESCENS,	Esterase
		EMBL:PF12537[PUTATIVE], PROTEIN (FRAGMENT)	
14966	·	CHITINASE CLASS IV (CHIV)	Chitinase
14969	1874	RNA POLYMERASE 24KDA SUBUNIT -LIKE PROTEIN	Polymerase
		RNA POLYMERASE SUBUNIT, ARABIDOPSIS	
		THALIANA,EMBL:AF019248	]
14985	1875	SUGAR-PHOSPHATE ISOMERASE - LIKE PROTEIN	Isomerase
		GUTQ/KPSF FAMILY SUGAR-P ISOMERASE,	
	i	CHLAMYDIA PNEUMONIAE, PIR:E72068	
14986	1876	PHOSPHATE TRANSPORT PROTEIN PHOSPHATE	Transporter
		TRANSPORTER, NICOTIANA TABACUM,	
		EMBL:AB020061	
14999	1877	VESICULAR TRANSPORTER UNC-47, CAENORHABDITIS	Transporter
		ELEGANS, AF031935[PUTATIVE]	-
15008	1878	PECTATE LYASE - LIKE PROTEIN PECTATE LYASE;	Lyase
		MUSA ACUMINATA, X92943	
15019	1879	PHOSPHATIDYLGLYCEROPHOSPHATE SYNTHASE -	Synthase
		LIKE PROTEIN PHOSPHATIDYLGLYCEROPHOSPHATE	
	i	SYNTHASE, PSEUDOMONAS FLUORESCENS,	
		EMBL:L29642	
15021	1880	PROTEIN PHOSPHATASE 2C - LIKE PROTEIN PROTEIN	Phosphatase
		PHOSPHATASE 2C HOMOLOG, MESEMBRYANTHEMUM	,
		CRYSTALLINUM, EMBL AF097667	
15028	1881	CHALCONE ISOMERASE	Isomerase
15034		LIPASE -LIKE PROTEIN MONOGLYCERIDE LIPASE,	
		MUS MUSCULUS, EMBL:MMMGLYLIP	
15035	1883	LIPASE -LIKE PROTEIN MONOGLYCERIDE LIPASE,	Linace
15055		MUS MUSCULUS, EMBL:MMMGLYLIP	Lipase
15041			Lugae
13041		-	Lyase
15040		PIR:T06728[PUTATIVE]	
15043		1	Phosphatase
	[	PHOSPHATASE - LIKE PROTEIN, RATTUS	}
		NORWEGICUS, SWISSPROT:DUS5_RAT	

PCT/EP01/09892

15045	1886	ALCOHOL DEHYDROGENASE HOMOLOG, TOMATO	Dehydrogenases
		PIR:S3950811BETA-HYDROXYSTEROID	
		DEHYDROGENASE (EC 1.1.1.146) 1 - MOUSE,	1
		PIR1:I56604[PUTATIVE] OXIDOREDUCTASE -LIKE	
		PROTEIN RIPENING-RELATED	1
15047	1887	BETA-KETOACYL-ACP REDUCTASE - LIKE PROTEIN	Reductase
		BETA-KETOACYL-ACP REDUCTASE, CUPHEA	
		LANCEOLATA, EMBL:X64566	
15059	1888	BETA-1,3-GLUCANASE - LIKE PROTEIN PROBABLE	Glycosylase
		BETA-1,3-GLUCANASE, TRITICUM AESTIVUM,	
		PIR:T06268	
15061	1889	SERINE/THREONINE-SPECIFIC PROTEIN KINASE -LIKE	1
		SERINE/THREONINE-SPECIFIC PROTEIN KINASE NAK,	
		ARABIDOPSIS THALIANA, PIR:S38326	
15086	1890	GLUCURONOSYL TRANSFERASE - LIKE PROTEIN	Transferases
		GLUCURONOSYL TRANSFERASE HOMOLOG,	
15087	1001	LYCOPERSICON ESCULENTUM, PIR:S39507	T
13087	1991	GLUCURONOSYL TRANSFERASE - LIKE PROTEIN GLUCURONOSYL TRANSFERASE HOMOLOG,	
		LYCOPERSICON ESCULENTUM, PIR:S39507	
15090	1892	PROLINE TRANSPORTER 2	Transporter
15094		BETA-1,3-GLUCANASE - LIKE PROTEIN BETA-1,3-	L
		GLUCANASE, ARABIDOPSIS THALIANA, PIR:S31906	
15096	1894		Phosphatase
15102	1005		
15103	1895	ANTHRANILATE SYNTHASE ALPHA-1 CHAIN - LIKE	-
		PROTEIN ANTHRANILATE SYNTHASE ALPHA	-
15111	1806	SUBUNIT, RUTA GRAVEOLENS, EMBL:RGANTSYNB RECEPTOR KINASE - LIKE PROTEIN RECEPTOR KINASE	Vinosa Protain
111161	1070	HOMOLOG CRINKLY4, MAIZE, PIR:T04108	kniase, flotein
15113	1897		Oxygenases
		PROTEIN LEUCOANTHOCYANIDIN DIOXYGENASE,	, ,
		APPLE TREE, PIR:S33144	
15116	1898	CELLULOSE SYNTHASE (CELA), AGROBACTERIUM	Synthase
	!	TUMEFACIENS, EMBL:ATCELABCE[PUTATIVE]	
LL		L	

15121	1899	PROTEIN KINASE SERINE/THREONINE-SPECIFIC	Kinase, Protein
		PROTEIN KINASE NPK15 - NICOTIANA TABACUM	
		PIR:S52578[PUTATIVE].	
15122	1900	MANDELONITRILE LYASE-LIKE PROTEIN ADHESION	Lyase
1		OF CALYX EDGES (ACE) - ARABIDOPSIS THALIANA	
15129	1901	ACETYL-COA CARBOXYLASE, BIOTIN CARBOXYL	Carboxylase
		CARRIER - ANABAENA SP., PIR:B53311[PUTATIVE]	
15136	1902	GLUTAMINE TRANSPORTER NEURONAL - RATTUS	Transporter
		NORVEGICUS, EMBL:AF075704[PUTATIVE]	:
15144	1903	PROTEIN KINASE NPK1-RELATED PROTEIN KINASE 1S	Kinase, Protein
		- ARABIDOPSIS THALIANA, EMBL:AB000797	
15148	1904	POLY(A) POLYMERASE - CANDIDA ALBICANS,	Polymerase
		EMBL:AB009394[PUTATIVE]	
15149	1905	N2,N2-DIMETHYLGUANOSINE TRNA	Transferases
		METHYLTRANSFERASES-LIKE PROTEIN SEVERAL	
		N2,N2-DIMETHYLGUANOSINE TRNA	
		METHYLTRANSFERASES	
15153	1906	LEUCINE-RICH RECEPTOR-LIKE PROTEIN KINASE -	Kinase, Protein
		MALUS DOMESTICA, EMBL:AF053127[PUTATIVE]	
15162	1907	QUINONE REDUCTASE-LIKE PROTEIN ZETA-	Reductase
		CRYSTALLIN / QUINONE REDUCTASE (NADPH) - MUS	
		MUSCULUS, PIR:A54932	·
15200			Dehydrogenases
		BACTERIA[PUTATIVE]	
15224			Polymerase
		POLYPEPTIDE (RPB25) - HOMO SAPIENS,	
		EMBL:J04965[PUTATIVE]	
15240			Glycosylase
15242			Glycosylase
15243	1	GLUCAN ENDO-1,3-BETA-D-GLUCOSIDASE-LIKE	Glycosylase
İ		PROTEIN GLUCAN ENDO-1,3-BETA-D-GLUCOSIDASE -	ł
		GLYCINE MAX, PIR:T07108	
15263	1		Protease
		RATTUS SP, EMBL: X93208[PUTATIVE]	
15264		PROTEASE-LIKE PROTEIN INSULIN-DEGRADING	Protease
		ENZYME - RATTUS NORVEGICUS, SWISSPROT:P35559	

15268	1915	ENDO-POLYGALACTURONASE	Glycosylase
15273	1916	ACETYLGLUTAMATE KINASE-LIKE PROTEIN	Kinase
		ACETYLGLUTAMATE KINASE - SYNECHOCYSTIS SP.,	
		PIR:S77509	
15279	1917	GALACTOSE OXIDASE PRECURSO, CLADOBOTRYUM	Oxidase
		DENDROIDES, PIR2:A38084[PUTATIVE]	
15281	1918	WALL-ASSOCIATED KINASE 1, ARABIDOPSIS	Kinase, Protein
		THALIANA, AJ009696[PUTATIVE]	
15283	1919	DNA-DIRECTED RNA POLYMERASE I 190K CHAIN -	
		LIKE PROTEIN DNA-DIRECTED RNA POLYMERASE(EC	
		2.7.7.6) I 190K CHAIN, SACCHAROMYCES CEREVISIAE,	
		PIR2:S67250	
15285	1920	CARBOXYL TERMINAL PROTEASE - LIKE PROTEIN	Protease
		CARBOXYL TERMINAL PROTEASE, NOSTOC	
1.50.05	1001	PUNCTIFORME, AF022823	,
15287	1921	WALL-ASSOCIATED KINASE 4, ARABIDOPSIS	Kinase, Protein
15000	1000	THALIANA,AJ009695	
15288	1922	WALL-ASSOCIATED KINASE 4, ARABIDOPSIS	Kinase, Protein
15280	1022	THALIANA, AJ009695	Vinesa Dustain
15289	1923	PROTEIN KINASE - LIKE PROTEIN S-RECEPTOR KINASE (EC 2.7.1) PRECURSOR, BRASSICA OLERACEA,	Kinase, Protein
		PIR1:S31429	
15290	1924	WALL-ASSOCIATED KINASE 1, ARABIDOPSIS	Kinase Protein
13290	1724	THALIANA, AJ009696[PUTATIVE]	ixinase, i rotem
15291	1925	WALL-ASSOCIATED KINASE 4, ARABIDOPSIS	Kinase Protein
13251	.,25	THALIANA, AJ009695[PUTATIVE]	12
15292	1926	WALL-ASSOCIATED KINASE I, ARABIDOPSIS	Kinase, Protein
		THALIANA, AJ009696	, , , , , , , , , , , , ,
15293	1927	WALL-ASSOCIATED KINASE 1, ARABIDOPSIS	Kinase, Protein
		THALIANA, AJ009696[PUTATIVE]	
15294	1928	WALL-ASSOCIATED KINASE 4, ARABIDOPSIS	Kinase, Protein
		THALIANA, AJ009695	
15296	1929	POLYGALACTURONASE (EC 3.2.1.15) PRECURSOR,	Glycosylase
		ERWINIA CAROTOVORA, PIR:S11773[PUTATIVE]	
15300	1930	RECEPTOR-LIKE PROTEIN KINASE SEVERAL	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASES	
L			

15305	1931	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE	Transferases
ļ		LIKE PROTEIN ANTHRANILATE	
		PHOSPHORIBOSYLTRANSFERASE - PISUM SATIVUM.	J
		PIR:T06460	
15327	1932	GLUCAN ENDO-1,3-BETA-GLUCOSIDASE PRECURSOR -	Glycosylase
		TRITICUM AESTIVUM, SWISSPROT:P52409[PUTATIVE]	
15366	1933		Transporter
		SACCHAROMYCES CEREVISIAE, PIR:S55178[PUTATIVE]	}
15378	1934	KETOL-ACID REDUCTOISOMERASE	Isomerase
15386	<u> </u>	SERINE/THREONINE-SPECIFIC PROTEIN KINASE -LIKE	Kinase, Protein
		  PROTEIN SERINE/THREONINE-SPECIFIC PROTEIN	i .
		KINASE NAK, ARABIDOPSIS THALIANA, PIR:S38326	
15391	1936	CITRATE SYNTHASE -LIKE PROTEIN CITRATE	Synthase
		SYNTHASE, CUCURBIT, PIR:S53007	
15392	1937	CITRATE SYNTHASE -LIKE PROTEIN CITRATE	Synthase
		SYNTHASE, CUCURBIT, PIR:S53007	
15393	1938	TYROSINE KINASE, DICTYOSTELIUM DISCOIDEUM,	Kinase, Protein
		PIR:A35670[PUTATIVE]	
15399	1939	RIBOSOMAL RNA APURINIC SITE SPECIFIC LYASE,	Lyase
		TRITICUM AESTIVUM, EMBL:AB032123[PUTATIVE]	
15416	1940	3-ISOPROPYLMALATE DEHYDRATASE-LIKE PROTEIN	Dehydratase
		(SMALL SUBUNIT) 3-ISOPROPYLMALATE	
		DEHYDRATASE, SMALL SUBUNIT - THERMOTOGA	
		MARITIMA, PIR:A72363	
15418	1941	PECTINESTERASE PRECURSOR-LIKE PROTEIN	Esterase
		PECTINESTERASE (EC 3.1.1.11) PRECURSOR - PISUM	
		SATIVUM, PIR:T06468	
15427	1942	CALLOSE SYNTHASE CATALYTIC SUBUNIT (CFL1) -	Synthase
		GOSSYPIUM HIRSUTUM, EMBL:AF085717[PUTATIVE]	
15428	1943	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE	Kinase, Protein
		PROTEIN KINASE (RKS2) - ARABIDOPSIS THALIANA,	
		EMBL:AF084036	
15431	1944	ABC TRANSPORTER-LIKE PROTEIN PUTATIVE MULTI	Transporter
		RESISTANCE PROTEIN MRP - ARABIDOPSIS THALIANA,	
		EMBL:ATMRPPROT	

INTERACTOR 1 - LYCOPERSICON ESCULENTUM, EMBL:U28007  15453 1946 TRANSPORTER-LIKE PROTEIN UDP-GALACTOSE Transporter TRANSPORT PROTEIN HOMOLOG SCHIZOSACCHAROMYCES POMBE, PIR:T43506  15459 1947 RECEPTOR KINASE HOMOLOG CRINKLY4 - ZEA MAYS, Kinase, Protein PIR:T04108[PUTATIVE]  15465 1948 FRUCTOKINASE-LIKE PROTEIN FRUCTOKINASE Kinase LYCOPERSICON ESCULENTUM, EMBL:LEUG2329  15487 1949 SERINE/THREONINE-SPECIFIC KINASE LECRKI Kinase, Protein PRECURSOR, LECTIN RECEPTOR-LIKE  15490 1950 RECEPTOR LECTIN KINASE LIKE PROTEIN Kinase, Protein (FRAGMENT) RECEPTOR LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15491 1951 RECEPTOR LECTIN KINASE 3 - Kinase, Protein LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15492 1952 RECEPTOR LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15500 1953 SERINE/THREONINE-PROTEIN KINASE CTR1 - Kinase, Protein ARABIDOPSIS THALIANA, EMBL:L08789[PUTATIVE]  15502 1954 POLYGALACTURONASE - PRUNUS PERSICA, PIR:S40123  15506 1955 DIHYDRODIPICOLINATE REDUCTASES REductase (DAPB)[PUTATIVE]  15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15602 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]	15452	1945	PROTEIN KINASE-LIKE PROTEIN PTO KINASE	Kinase, Protein
15453 1946 TRANSPORTER-LIKE PROTEIN UDP-GALACTOSE TRANSPORT PROTEIN HOMOLOG - SCHIZOSACCHAROMYCES POMBE, PIR:T43506  15459 1947 RECEPTOR KINASE HOMOLOG CRINKLY4 - ZEA MAYS, Kinase, Protein PIR:T04108[PUTATIVE]  15465 1948 FRUCTOKINASE-LIKE PROTEIN FRUCTOKINASE - Kinase LYCOPERSICON ESCULENTUM, EMBL:LEU62329  15487 1949 SERINE/THREONINE-SPECIFIC KINASE LECRKI Kinase, Protein PRECURSOR, LECTIN RECEPTOR-LIKE  15490 1950 RECEPTOR LECTIN KINASE-LIKE PROTEIN Kinase, Protein (FRAGMENT) RECEPTOR LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15491 1951 RECEPTOR LECTIN KINASE 3 - Kinase, Protein LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15492 1952 RECEPTOR LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15500 1953 SERINE/THREONINE-PROTEIN KINASE CTR1 - Kinase, Protein ARABIDOPSIS THALIANA, EMBL:L08789[PUTATIVE]  15500 1954 POLYGALACTURONASE-LIKE PROTEIN Glycosylase POLYGALACTURONASE - PRUNUS PERSICA, PIR:S40123  15506 1955 DIHYDRODIPICOLINATE REDUCTASES Reductase (DAPB)[PUTATIVE]  15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15500 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15600 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15600 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease				
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LECTIN KINASE 3 - ARABIDOPSIS THALIANA, EMBL:U93161  15500 1953 SERINE/THREONINE-PROTEIN KINASE CTR1 - Kinase, Protein ARABIDOPSIS THALIANA, EMBL:L08789[PUTATIVE]  15502 1954 POLYGALACTURONASE-LIKE PROTEIN Glycosylase POLYGALACTURONASE - PRUNUS PERSICA, PIR:S40123  15506 1955 DIHYDRODIPICOLINATE REDUCTASES Reductase (DAPB)[PUTATIVE]  15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase  15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15491	1951	RECEPTOR LECTIN KINASE 3	Kinase, Protein
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15506 1955 DIHYDRODIPICOLINATE REDUCTASES Reductase (DAPB)[PUTATIVE]  15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase 15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15502	1954	POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
(DAPB)[PUTATIVE]  15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase 15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease			POLYGALACTURONASE - PRUNUS PERSICA, PIR:S40123	
15579 1956 PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE Transferases PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase 15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15506	1955	DIHYDRODIPICOLINATE REDUCTASES	Reductase
PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE, SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase 15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease			(DAPB)[PUTATIVE]	
SYNECHOCYSTIS SP., PIR:S77254  15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase 15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15579	1956	PHOSPHATIDATE CYTIDYLYLTRANSFERASE - LIKE	Transferases
15590 1957 PECTINESTERASE - LIKE PROTEIN PECTINESTERASE, Esterase SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase  15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease			PROTEIN PHOSPHATIDATE CYTIDYLYLTRANSFERASE,	
SWEET ORANGE, PIR:T10494  15605 1958 DIHYDRODIPICOLINATE SYNTHASE PRECURSOR Synthase  15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease			SYNECHOCYSTIS SP., PIR:S77254	
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15612 1959 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease			SWEET ORANGE, PIR:T10494	
EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]  15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15605	1958	DIHYDRODIPICOLINATE SYNTHASE PRECURSOR	Synthase
15620 1960 APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA, Nuclease	15612	1959	APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA,	Nuclease
			EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]	
EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]	15620	1960	APURINIC ENDONUCLEASE - ARABIDOPSIS THALIANA,	Nuclease
			EMBL:X76912 (AT C-TERMINUS)[PUTATIVE]	

- 159 -

15665	1000	ANIMATE AND DATE OF THE PROPERTY OF THE PROPER	Tes
15647	1961	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE-	
		LIKE PROTEIN ANTHRANILATE	
		PHOSPHORIBOSYLTRANSFERASE (FRAGMENT) -	
		PISUM SATIVUM, PIR:T06460	
15657	1962	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE-	Oxidase
		LIKE PROTEIN 1-AMINOCYCLOPROPANE-1-	
		CARBOXYLATE OXIDASE HOMOLOG - ARABIDOPSIS	
		THALIANA, PIR:S59548	
15658	1963	PROTEIN KINASE HOMOLOG F4118.11 - ARABIDOPSIS	Kinase, Protein
		THALIANA, PIR:T02456[PUTATIVE]	
15666	1964	POLYGALACTURONASE - ZEA MAYS,	Glycosylase
		EMBL:ZMPGAL3[PUTATIVE]	
15670	1965	3-METHYL-2-OXOBUTANOATE HYDROXY-METHYL-	Transferases
		TRANSFERASE-LIKE PROTEIN KETOPANTOATE	
		HYDROXYMETHYLTRANSFERASE - EMERICELLA	
		NIDULANS, EMBL:AF134703	
15671	1966	PROLYL AMINOPEPTIDASE-LIKE PROTEIN PROLYL	Protease
		AMINOPEPTIDASE - AEROMONAS SOBRIA, PIR:JC4184	
15689	1967	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE -	Transferases
		PISUM SATIVUM, EMBL:D86180[PUTATIVE]	
15698	1968	BETA-1,3-GLUCANASE PRECURSOR - ORYZA SATIVA,	Glycosylase
		EMBL:U72255[PUTATIVE]	
15716	1969	O-METHYLTRANSFERASE - DIFFERENT	Transferases
		SPECIES[PUTATIVE]	
15723	1970	PECTINACETYLESTERASE PRECURSOR-LIKE PROTEIN	Esterase
		PECTINACETYLESTERASE PRECURSOR - VIGNA	
		RADIATA, EMBL:X99348	
15728	1971	POLYGALACTURONASE - LYCOPERSICON	Glycosylase
		ESCULENTUM, EMBL:AF118567[PUTATIVE]	
15730	1972	ISOPENICILLIN N EPIMERASE - STREPTOMYCES	Epimerase
		CLAVULIGERUS, EMBL:M32324[PUTATIVE]	
15733	1973	10-DEACETYLBACCATIN III-10-O-ACETYL	Transferases
	1	TRANSFERASE - TAXUS CUSPIDATA,	
		EMBL:AF193765[PUTATIVE]	-
15734	1974	PECTINESTERASE-LIKE PROTEIN PECTINESTERASE -	Esterase
	i	BRASSICA RAPA, EMBL:L48178	Ì

15735	1975	PECTIN METHYLESTERASE - ARABIDOPSIS THALIANA	, Esterase
		EMBL:AJ250430[PUTATIVE]	
15768	1976	protein DISULFIDE-ISOMERASE - ZEA MAYS	Isomerase
		PIR:S69181[PUTATIVE]	
15782	1977	MITOCHONDRIAL CARRIER PROTEIN AT2G47490 -	Transporter
		ARABIDOPSIS THALIANA, EMBL:AC002535[PUTATIVE]	
15783	1978	GLYCOSYL TRANSFERASE LGTC - NEISSERIA	Transferases
		GONORRHOEAE, EMBL:AF208062[PUTATIVE]	
15787	1979	ABC TRANSPORTER-LIKE PROTEIN GLUTATHIONE-	Transporter
		CONJUGATE TRANSPORTER ATMRP4 - ARABIDOPSIS	
		THALIANA, EMBL:AJ002584	}
15788	1980	BETA-D-GLUCAN EXOHYDROLASE-LIKE PROTEIN	Hydrolase
		EXHYDROLASE II - ZEA MAYS, EMBL:AF064707	
15789	1981	ALPHA GALACTOSYLTRANSFERASE-LIKE PROTEIN	Transferases
		ALPHA GALACTOSYLTRANSFERASE - TRIGONELLA	
		FOENUM-GRAECUM, EMBL:TFO245478	
15793	1982	GLUTATHIONE TRANSFERASE III-LIKE PROTEIN	Transferases
		GLUTATHIONE TRANSFERASE III(B) - ZEA MAYS,	
		EMBL:AJ010296	
15799	1983	PECTINESTERASE HOMOLOG - PINUS RADIATA,	Esterase
		PIR:T08112[PUTATIVE]	
15803		LYSOPHOSPHOLIPASE HOMOLOG - ORYZA SATIVA,	Lipase
		PIR:T02661[PUTATIVE]	
15805			Channel
		HORDEUM VULGARE, EMBL:AJ011921[PUTATIVE]	
15828	1986	TRNA ISOPENTENYL TRANSFERASE -LIKE PROTEIN	Transferases
] ]		TRNA ISOPENTENYL TRANSFERASE, ARABIDOPSIS	
1-0.15		THALIANA, EMBL:AF109376	
15845	ł	SERINE/THREONINE-PROTEIN KINASE NEK4 - MUS	Kinase, Protein
1.50.10		MUSCULUS, EMBL:AF099067[PUTATIVE]	
15849	i	PROTEIN PHOSPHATASE 2C - RATTUS NORVEGICUS,	Phosphatase
15050		EMBL:AF095927[PUTATIVE]	
15850	1989	MAP KINASE [PUTATIVE]	Kinase, Protein
15851	1990	PROTEIN PHOSPHATASE 2C - RATTUS NORVEGICUS,	Phosphatase
		EMBL:AF095927[PUTATIVE]	

15861	1991	CYTOKININ OXIDASE -LIKE PROTEIN CYTOKININ	Oxidase
		OXIDASE, ZEA MAYS, EMBL:ZMY18377	
15873	1992	CHALCONE SYNTHASE SIMILAR TO PLANT CHALCONE	Synthase
		AND STILBENE SYNTHASES[PUTATIVE]	
15880	1993	NUCLEOTIDE SUGAR EPIMERASE [PUTATIVE]	Epimerase
15888	1994	PECTINESTERASE [PUTATIVE]	Esterase
15892	1995	CUCUMISIN PROTEASE [PUTATIVE]	Protease
15893	1996	PHOSPHOLIPASE D-LIKE PROTEIN	Lipase
15897	1997	LIGHT REPRESSIBLE RECEPTOR PROTEIN KINASE	Kinase, Protein
		(PID:E242366), SIMILAR TO A. THALIANA	
15903	1998	S-DOMAIN RECEPTOR-LIKE PROTEIN KINASE, ZEA	Kinase, Protein
		MAYS, SIMILARITY TO	
15918	1999	BETA-AMYLASE SIMILAR TO THE FAMILY OF	Hydrolase
		GLYCOSYL HYDROLASES[PUTATIVE]	
15921	2000	ACETYL COA THIOESTERASE [PUTATIVE]	Esterase
15925	2001	DTDP-6-DEOXY-L-MANNOSE-DEHYDROGENASE	Dehydrogenases
		[PUTATIVE]	
15932	2002	POTASSIUM/H+ ANTIPORTER [PUTATIVE]	Transporter
15939	2003	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE	Transferases
		[PUTATIVE]	
15970	2004	CYCLIC NUCLEOTIDE GATED CHANNEL (CNGC4) LIKE	Channel
		PROTEIN ARABIDOPSIS THALIANA CYCLIC	
		NUCLEOTIDE GATED CHANNEL (CNGC4),PID:G4378659	
15976	2005	FLAVONOL GLUCOSYLTRANSFERASE SIMILAR TO	Transferases
		MANIHOT ESCULENTA FLAVONOL 3-0-	
		GLUCOSYLTRANSFERASE 5, GENBANK ACCESSION	
		NUMBER Q40287[PUTATIVE]	
15982	2006	ACETYLTRANSFERASE SIMILAR TO DIGITALIS	Transferases
		LANATA LANATOSIDE 15'-O-ACETYLTRANSFERASE,	
		GENBANK ACCESSION NUMBER AJ011567[PUTATIVE]	
16002	2007	PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16011	2008	CALCIUM-REGULATED PROTEIN PHOSPHATASE	Phosphatase
		SIMILAR TO N. CRASSA CALCINEURIN CALCIUM-	
		REGULATED PROTEIN PHOSPHATASE, GENBANK	
		ACCESSION NUMBER P87072[PUTATIVE]	
16016	2009	WATER CHANNEL PROTEIN [PUTATIVE]	Channel

- 162 -

16017	2010	INORGANIC PHOSPHATASE [PUTATIVE]	Phosphatase
16022	2011	REVERSE TRANSCRIPTASE-LIKE PROTEIN [PUTATIVE]	Transcriptase
16037	2012	PROTOPORPHYRINOGEN OXIDASE	Oxidase
16038	2013	CHITINASE SIMILAR TO PEANUT TYPE II CHITINASE,	Chitinase
		GENBANK ACCESSION NUMBER X82329, E.C.	
		3.2.1.14[PUTATIVE]	
16048	2014	SECA-TYPE CHLOROPLAST PROTEIN TRANSPORT	Transporter
		FACTOR [PUTATIVE]	
16049	2015	PROTEIN TRANSPORT FACTOR [PUTATIVE]	Transporter
16052	2016	POTASSIUM CHANNEL [PUTATIVE]	Channel
16065	2017	RAFFINOSE SYNTHASE OR SEED IMBIBITION PROTEIN	Synthase
		SIMILAR TO CUCUMBER RAFFINOSE SYNTHASE,	
		GENBANK ACCESSION NUMBER AF073744[PUTATIVE]	
16069	2018	NAK-LIKE SER/THR PROTEIN KINASE SIMILAR TO A.	Kinase, Protein
		THALIANA NAK SER/THR PROTEIN KINASE, GENBANK	
		ACCESSION NUMBER P43293[PUTATIVE]	
16081	2019	GLYCOSYL TRANSFERASE SIMILAR TO LGTC OF	Transferases
		NEISSERIA SP., GENBANK ACCESSION NUMBER	
		U14554, SIMILAR TO LGTC, GENBANK ACCESSION	
		NUMBER U65788	
16094	2020	GTP PYROPHOSPHOKINASE SIMILAR TO BACTERIAL	Kinase
		GTP PYROPHOSPHOKINASES (RELA) SIMILAR TO B.	
		SUBTILIS RELA (EC 2.7.6.5), GENBANK ACCESSION	
		NUMBER 2635224 CARBOXYL REGION OF PROTEIN	
		NOT DEFINED[PUTATIVE]	
16096		SUCROSE SYNTHETASE SIMILAR TO SEVERAL PLANT	Synthase
		SUGAR SYNTHETASES SIMILAR TO P. SATIVUM	•
	j	SECOND SUGAR SYNTHETASE, GENBANK ACCESSION	
	1	NUMBER AJ001071 SIMILAR TO BEET SUCROSE	
		SYNTHETASE (EC 2.4.1.13), GENBANK ACCESSION	
1,000		NUMBER S71494[PUTATIVE]	D
16098	1	PECTINESTERASE SIMILAR TO SEVERAL PLANT	Esterase
16100		PECTINESTERASES[PUTATIVE]	r
16100	1	PECTINESTERASE - LIKE PROTEIN SIMILAR TO	Esterase
		BACTERIAL AND FUNGI PECTINESTERASES	

16101	2024	PECTINESTERASE, SIMILAR TO	Esterase
16118	2025	GLYCOSYLTRANSFERASE SIMILAR TO A. THALIANA	Transferases
		PROTEIN T20K9.11, GENBANK ACCESSION NUMBER	
		3445207[PUTATIVE]	
16120	2026	GLUTATHIONE S TRANSFERASE, ATPM24.1	Transferases
16129	2027	TRYPTOPHAN SYNTHASE ALPHA 1-LIKE PROTEIN	Synthase
		SIMILAR TO A. THALIANA TRYPTOPHAN SYNTHASE	
		ALPHA CHAIN (EC 4.2.1.20), GENBANK ACCESSION	
		NUMBER U18993	
16131	2028	SERINE/THREONINE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16137	2029	Glutamate N-METHYL-D-ASPARTATE RECEPTOR;	Receptor
	•	GLUTAMATE-/ASPARTATE-BINDING PEPTIDE SIMILAR	
		TO RAT N-METHYL-D-ASPARTATE RECEPTOR	
		GLUTAMATE-BINDING CHAIN, GENBANK ACCESSION	
		NUMBER S19586[PUTATIVE]	
16146	2030	ENT-KAURENE SYNTHETASE A - LIKE PROTEIN	Synthase
16173	2031	OXIDOREDUCTASE SIMILAR TO A. THALIANA	Reductase
		PUTATIVE PROTEIN F21P8.230, GENBANK ACCESSION	
		NUMBER 3445238[PUTATIVE]	
16174	2032	GIBBERELLIN 20-OXIDASE, GENBANK ACCESSION	Oxidase
		NUMBER U70530[PUTATIVE], OXIDOREDUCTASE	
		SIMILAR TO P. VULGARIS	
16175		-	Reductase
16182	2034	ALCOHOL DEHYDROGENASE SIMILAR TO PLANT	Dehydrogenases
		SHORT CHAIN ALCOHOL	
		DEHYDROGENASE[PUTATIVE]	
16189		XYLOGLUCAN ENDOTRANSGLYCOSYLASE	Glycosylase
		[PUTATIVE]	
16191	2036	RECEPTOR KINASE [PUTATIVE]	Kinase, Protein
16194	2037	PROTEIN PHOSPHATASE REGULATORY SUBUNIT	Phosphatase
		[PUTATIVE]	
16202	2038	GLYCOSYLATION ENZYME SIMILAR TO B. TAURUS	Transferases
		CORE2-GLCNAC-TRANSFERASE, GENBANK	1
		ACCESSION NUMBER U41320[PUTATIVE]	

16207	2039	LRR RECEPTOR-LIKE PROTEIN KINASE SIMILAR TO Z.	Kinase, Protein
		MAYS LEUCINE-RICH REPEAT TRANSMEMBRANE	
		PROTEIN KINASE LRRTPK 1, GENBANK ACCESSION	
		NUMBER AF023164[PUTATIVE]	
16221	2040	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
16223	2041	GLUCAN SYNTHASE COMPONENT SIMILAR TO 1,3-	Synthase
		BETA GLUCAN SYNTHASE[PUTATIVE]	
16229	2042	HYDROLASE [PUTATIVE]	Hydrolase
16261	2043	PECTINESTERASE [PUTATIVE]	Esterase
16268	2044	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
16272	2045	PHOSPHOFRUCTOKINASE BETA SUBUNIT [PUTATIVE]	Kinase
16280	2046	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
16290	2047	DISEASE RESISTANCE PROTEIN SIMILAR TO	Kinase, Protein
		RECEPTOR PROTEIN KINASES[PUTATIVE]	
16317	2048	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16318	2049	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16319	2050	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16322	2051	RECEPTOR-LIKE PROTEIN KINASE [PUTATIVE]	Kinase, Protein
16325	2052	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
16329	2053	5-ADENYLYLSULFATE REDUCTASE	Reductase
16345	2054	ABC TRANSPORTER SIMILAR TO GUILLARDIA THETA	Transporter
		ABC TRANSPORTER, GENBANK ACCESSION NUMBER	
		AF041468[PUTATIVE]	
16353	2055	POTASSIUM TRANSPORTER SIMILAR TO A. THALIANA	Transporter
		K+ ANTIPORTER KEAI, GENBANK ACCESSION	
		NUMBER AF003382[PUTATIVE]	
16355	2056	PHOSPHATIDYLGLYCEROTRANSFERASE SIMILAR TO	Transferases
		CDP-DIACYLGLYCEROL-GLYCEROL-3-PHOSPHATE 3-	1
		PHOSPHATIDYLTRANSFERASE OF SYNECHOCYSTIS	
		SP. GENBANK ACCESSION NUMBER D90914[PUTATIVE]	
16379	2057	EQUILIBRATIVE NUCLEOSIDE TRANSPORTER 1, HOMO	Transporter
		SAPIENS, GB:NP_004946[PUTATIVE]	

16380	2058	EQUILIBRATIVE NUCLEOSIDE TRANSPORTER 1, HOMO	Transporter
10300	2030	SAPIENS, GB:NP_004946[PUTATIVE]	Transporter
1.6201	2050		
16381	2059	EQUILIBRATIVE NUCLEOSIDE TRANSPORTER 1, HOMO	Transporter
		SAPIENS, GB:NP_004946[PUTATIVE]	
16382	2060	EQUILIBRATIVE NUCLEOSIDE TRANSPORTER 1, HOMO	Transporter
		SAPIENS, GB:NP_004946[PUTATIVE]	
16388	2061	RECEPTOR PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
1		RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1,	
		PHAESOLUS VULGARIS, GB:AAD21872	
16404	2062	GAG-PROTEASE POLYPROTEIN, GLYCINE MAX.,	Protease
<u> </u>		GB:AAC18777[PUTATIVE]	
16406	2063	AAA FAMILY ATPASE BCS1P MITOCHONDRIAL,	ATPase
	]	SACCHAROMYCES CEREVISIAE, SWISS	,
		PROT:P32839[PUTATIVE]	
16407	2064	FERREDOXINNADP+ REDUCTASE - LIKE PROTEIN	Reductase
		FERREDOXINNADP+ REDUCTASE, PISUM SATIVUM,	
		PIR:T06773	
16411	2065	GLUCAN ENDO-1,3-BETA-GLUCOSIDASE PRECURSOR,	Glycosylase
		TRITICUM AESTIVUM, U30323[PUTATIVE]	
16479	2066	REVERSE TRANSCRIPTASE [PUTATIVE]	Transcriptase
		-	-
16501	2067	GLUCOSYLTRANSFERASE [PUTATIVE]	Transferases
16516	2068	XYLAN ENDOHYDROLASE [PUTATIVE]	Hydrolase
16517	2069	INOSITOL 1,3,4-TRISPHOSPHATE 5/6-KINASE-LIKE	Kinase
		PROTEIN INOSITOL 1,3,4-TRISPHOSPHATE 5/6-KINASE	
		(EC 2.7) - ARABIDOPSIS THALIANA, PIR2:JC5401	
16526	2070	PROTEIN PHOSPHATASE 2C - MEDICAGO SATIVA,	Phosphatase
		PID:E305311[PUTATIVE]	
16539	2071	ASCORBATE PEROXIDASE STROMAL	Oxidase
16577	2072	PEROXIDASE C2 PRECURSOR LIKE PROTEIN	Oxidase
·		PEROXIDASE (EC 1.11.1.7) C2 PRECURSOR -	
		ARMORACIA RUSTICANA,PID:D1014846	
16578	2073	PEROXIDASE C2 PRECURSOR LIKE PROTEIN	Oxidase
		PEROXIDASE (EC 1.11.1.7) C2 PRECURSOR -	
		ARMORACIA RUSTICANA,PID:D1014846	

16583	2074	REVERSE TRANSCRIPTASE OF ARABIDOPSIS	Transcriptore
10505	2074	THALIANA[PUTATIVE]	Transcriptase
16586	2075	ARGINASE SIMILAR TO ARGINASES (PFAM: PF00491,	AMIDASE
10500	2075	SCORE=353.2, E=1.4E-119, N=1)[PUTATIVE]	AMIDASE
16589	2076	ARGINASE	ARGINASE.
16592		PROTEIN DISULFIDE ISOMERASE [PUTATIVE]	Isomerase
16600		ASCORBATE PEROXIDASE - SPINACIA	
10000	2076	OLERACEA,PIR2:S66265[PUTATIVE]	Oxidase
16601	2079	ISOAMYLASE-LIKE PROTEIN PROBABLE ISOAMYLASE	Glycorylasa
10001	2017	(EC 3.2.1.68) SU1 - ZEA MAYS,PIR2:T01321	Glycosylasc
16608	2080	BETA-1,3-GLUCANASE-LIKE PROTEIN PROBABLE	Glycosylase
10000	2000	BETA-1,3-GLUCANASE - TRITICUM	Ciyeosylase
		AESTIVUM,PIR2:T06268	
16648	2081	ANTHOCYANIN RHAMNOSYLTRANSFERASE -PETUNIA	Transferases
		X HYBRIDA, PID:G454253[PUTATIVE]	
16650	2082	PHOSPHONOPYRUVATE DECARBOXYLASE (EC 4.1.1) -	Decarboxylase
		METHANOBACTERIUM	
		THERMOAUTOTROPHICUM,PID:G2622714[PUTATIVE]	
16700	2083	11BETA-HYDROXYSTEROID DEHYDROGENASE (EC	Dehydrogenases
		1.1.1.146) 1 - MOUSE, PIR1:I56604[PUTATIVE]	
16703	2084	LIPASE-LIKE PROTEIN MONOGLYCERIDE LIPASE -	Lipase
		MUS MUSCULUS, PID:E1184892	
16705	2085	DNA-DIRECTED RNA POLYMERASE DNA-DIRECTED	Polymerase
		RNA POLYMERASE (EC 2.7.7.6) II LARGESTCHAIN -	
		MOUSE, PIR2:A28490[PUTATIVE]	
16710	1	SUCROSE-PHOSPHATE SYNTHASE - LIKE PROTEIN	Synthase
	ſ	SUCROSE-PHOSPHATE SYNTHASE, ZEA MAYS,	
		PIR2:JQ1329	
16724	- 1	FRUCTOKINASE - LIKE PROTEIN FRUCTOKINASE,	Kinase
16770		LYCOPERSICON ESCULENTUM, GB:U62329	
16729		POTASSIUM UPTAKE TRANSPORTER - LIKE PROTEIN	Transporter
	i	POTASSIUM TRANSPORT PROTEIN TRK1,	
16727		SACCHAROMYCES CEREVISIAE, PIR2:JU0466	V: P
16737	J	CLV1 RECEPTOR KINASE, ARABIDOPSIS THALIANA,	Kinase, Protein
		GB:U96879[PUTATIVE]	

- 167 -

16747	2090	FLAVANONE 3-BETA-HYDROXYLASE NARINGENIN 3-	Oxygenases
		DIOXYGENASE (EC 1.14.11.9) - CHINA ASTER	,
		PIR2:S32147[PUTATIVE]	,
16748	2091	FE(II)/ASCORBATE OXIDASE SRG1 PROTEIN	Oxidase
		ARABIDOPSIS THALIANA, PIR2:S44261[PUTATIVE]	
16749	2092	SUBTILISIN-LIKE PROTEASE SUBTILISIN-LIKE	Protease
	4	PROTEASE - LYCOPERSICON ESCULENTUM (TOMATO),	
]		PID:E1299610[PUTATIVE]	
16750	2093	SUBTILISIN-LIKE PROTEASE SUBTILISIN-LIKE	Protease
		PROTEASE - LYCOPERSICON ESCULENTUM,	
		PID:E1299610[PUTATIVE]	
16751	2094	SUBTILISIN-LIKE PROTEASE SUBTILISIN-LIKE	Protease
		PROTEASE - LYCOPERSICON	
		ESCULENTUM,PID:E1299610[PUTATIVE]	
16752	2095	SUBTILISIN-LIKE PROTEASE SUBTILISIN-LIKE	Protease
		PROTEASE - LYCOPERSICON	
		ESCULENTUM,PID:E1299610[PUTATIVE]	
16753	2096	SUBTILISIN-LIKE PROTEASE -LIKE PROTEIN	Protease
		SUBTILISIN-LIKE PROTEASE AIR3 -ARABIDOPSIS	
		THALIANA,PID:G3695019	
16773	2097	2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID	
		ALDOLASES FUNCTIONAL CATALOG	
		ID=01.05[PUTATIVE]	
16775	2098		Transporter
		ALBICANS OPTI, GENBANK ACCESSION NUMBER	
		U60973 FUNCTIONAL CATALOG ID=07.99[PUTATIVE]	
16790		RNA POLYMERASE II TRANSCRIPTION COFACTOR P15,	Polymerase
		HOMO SAPIENS, PIR2:A54670[PUTATIVE]	
16802			Phosphatase
		3.1.3.16) 2C - ARABIDOPSIS THALIANA,	
		PIR2:S55457[PUTATIVE]	
16827		PEROXIDASE ATP19A	Oxidase
16842			Transporter
		NIGRUM,PID:E1313696[PUTATIVE]	

- 168 -

16844	2103	SERINE/THREONINE KINASE-LIKE PROTEIN Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE RLK3 -
		ARABIDOPSIS THALIANA,PID:E1363211
16845	2104	SERINE/THRÉONINE KINASE-LIKE PROTEIN Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE RLK3 -
		ARABIDOPSIS THALIANA,PID:E136321
16846	2105	SERINE/THREONINE KINASE-LIKE PROTEIN Kinase, Protein
1		RECEPTOR-LIKE PROTEIN KINASE RLK3 -
		ARABIDOPSIS THALIANA,PID:E136321
16847	2106	SERINE/THREONINE KINASE-LIKE PROTEIN Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE RLK3 -
		ARABIDOPSIS THALIANA,PID:E136321
16848	2107	SERINE/THREONINE KINASE-LIKE PROTEIN Kinase, Protein
		(FRAGMENT) RECEPTOR-LIKE PROTEIN KINASE RKS1 -
		ARABIDOPSIS THALIANA,PID:G4008008
16850	2108	RECEPTOR-LIKE PROTEIN KINASE RLK3 - Kinase, Protein
		ARABIDOPSIS THALIANA,PID:E136321[PUTATIVE]
16851	2109	SERINE/THREONINE KINASE-LIKE PROTEIN Kinase, Protein
		(FRAGMENT) RECEPTOR-LIKE PROTEIN KINASE RLK3 -
		ARABIDOPSIS THALIANA,PID:E136321
16855	2110	BETA-PHOSPHOGLUCOMUTASE - LACTOBACILLUS Mutase
		SANFRANCISCO,PID:E1331347[PUTATIVE]
16859	2111	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE Transferases
		PHOSPHORIBOSYLANTHRANILATE TRANSFERASE,
		PISUM SATIVUM, D86180[PUTATIVE]
16869	2112	REVERSE TRANSCRIPTASE /TRANSPOSON, Transcriptase
		ARABIDOPSIS THALIANA[PUTATIVE]
16881	2113	PHOSPHOLIPASE D-Lipase D-Lipase
	•	GAMMA - ARABIDOPSIS
	i	THALIANA,PID:G2653885[PUTATIVE]
16882	2114	PHOSPHOLIPASE D-GAMMA PHOSPHOLIPASE D-Lipase
		GAMMA - ARABIDOPSIS
		THALIANA,PID:G2653885[PUTATIVE]
16883	2115	PHOSPHOLIPASE D-Lipase
		GAMMA - ARABIDOPSIS
		THALIANA,PID:G2653885[PUTATIVE]

16887	2116	PROTEIN KINASE - LIKE PROTEIN KI DOMAINI	Kinase, Protein
		INTERACTING KINASE 1 -ZEA MAYS,PIR2:T02053	
16888	2117	KI DOMAIN INTERACTING KINASE 1 -LIKE PROTEIN KI	Kinase, Protein
		DOMAIN INTERACTING KINASE 1 - ZEA	·
		MAYS,PID:G2735017	
16901	2118	TRANSPORT PROTEIN NA(+) DEPENDENT T	Transporter
		TRANSPORTER (SBF FAMILY) - AQUIFEX AEOLICUS,	
		PIR2:E70482[PUTATIVE]	
16923	2119	NUCLEOTIDE SUGAR EPIMERASE -LIKE PROTEIN	Epimerase
		NUCLEOTIDE SUGAR EPIMERASE -VIBRIO	
		VULNIFICUS,PID:G3093975	
16925	2120	COPPER AMINE OXIDASE LIKE PROTEIN (FRAGMENTI)	Oxidase
		COPPER AMINE OXIDASE - CICER	
		ARIETINUM,PID:E1335964	
16926	2121	COPPER AMINE OXIDASE LIKE PROTEIN (FRAGMENT2)	Oxidase
		COPPER AMINE OXIDASE - CICER	
		ARIETINUM,PID:E1335964	
16937	2122	PECTINESTERASE - CITRUS E	sterase
		SINENSIS,PID:G2098711[PUTATIVE]	<u>-</u>
16941	2123	TREHALOSE-6-PHOSPHATE PHOSPHATASE (ATTPPA)	hosphatase
		TREHALOSE-6-PHOSPHATE PHOSPHATASE -	
16000		ARABIDOPSIS THALIANA, PID:G2944178[PUTATIVE]	
16980	2124	HYDROLASE-LIKE PROTEIN 2-HYDROXY-6-H	lydrolase
[ [		OXOHEPTA-2,4-DIENOATE HYDROLASE (EC 3.7)	
16008	2126	SYNECHOCYSTIS SP., PIR2:S77427  OXIDOREDUCTASE - STREPTOMYCES R	
16998			eductase
17002		LIVIDANS,PID:G3293547[PUTATIVE]	
17002		OLEOYL-[ACYL-CARRIER-PROTEIN] HYDROLASE-LIKE H PROTEIN OLEOYL-[ACYL-CARRIER-PROTEIN]	ydrolase
[ [		]	
17005		HYDROLASE - BRASSICA NAPUS, PIR2:S40407  ENDOXYLOGLUCAN TRANSFERASE-LIKE PROTEIN TO	ranafaras
1,003		XYLOGLUCAN IRANSFERASE-LIKE PROTEIN II XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE (EC	iansierases
		3.2.1) - ARABIDOPSIS THALIANA, PIR2: C49539	
17006		ENDOXYLOGLUCAN TRANSFERASE-LIKE PROTEIN TI	rancferacec
17000		XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE (EC	141151014505
		3.2.1) - ARABIDOPSIS THALIANA,PIR2:C49539	!
		7.5.1. / INCLUDED OF STREET, INC. C47337	

17015	2129	SHORT-CHAIN ALCOHOL DEHYDROGENASE LIKE	Dehydrogenase
		PROTEIN SHORT-CHAIN ALCOHOL DEHYDROGENASE	-
		PICEA ABIES, PIR2:S34678	
17018	2130	PECTATE LYASE LIKE PROTEIN PECTATE LYASE	- Lyase
		FRAGARIA X ANANASSA,PID:G2435395	
17023	2131	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE	Oxygenases
		FORMING) (EC1.14.13.8) -ORYCTOLAGUS	<b>;</b>
		CUNICULUS,PID:G164989[PUTATIVE]	
17027	2132	TERPENE SYNTHASE TS1, ARABIDOPSIS THALIANA	Synthase
		Y11188[PUTATIVE]	
17038	2133	CYCLIC BETA-1-3-GLUCAN SYNTHASE,	Synthase
		BRADYRHIZOBIUM JAPONICUM, AF047687[PUTATIVE]	
17039	2134	POTASSIUM TRANSPORTER - LIKE PROTEIN	Transporter
		POTASSIUM TRANSPORTER ATKTIP, ARABIDOPSIS	
		THALIANA,AF012656	
17040	2135	3-ISOPROPYLMALATE DEHYDRATASES/ACONITATE	Dehydratase
		HYDRATASES[PUTATIVE]	
17057	2136	BETA 1,3-GLUCANASE, TRITICUM AESTIVUM,	Glycosylase
		U30323[PUTATIVE]	
17063	2137	ISOFLAVONE REDUCTASE-LIKE PROTEIN	Reductase
		ISOFLAVONE REDUCTASE-LIKE PROTEIN, LUPINUS	
		ALBUS, GB:U48590	
17067	2138	PURPLE ACID PHOSPHATASE IPOMOEA BATATAS,	Phosphatase
		AJ006224[PUTATIVE]	
17068	2139	PECTATE LYASE A11 (FRAGMENT) [PUTATIVE]	Lyase
17073	2140	POLYGALACTURONASE POLYGALACTURONASE, ZEA	Glycosylase
		MAYS, PIR2:S30067[PUTATIVE]	
17106	2141	GLUCOSYLTRANSFERASE LIKE PROTEIN	Transferases
17110	2142	XYLOGLUCAN ENDOTRANSGLYCOSYLASE-RELATED	Glycosylase
		PROTEIN XTR-7	-
17118	2143	PHYTOENE DESATURASE, PHYTOENE	Desaturases
		DEHYDROGENASE PRECURSOR	
17140	2144	CARNITINE RACEMASE LIKE PROTEIN	Epimerase
17141	2145	CARNITINE RACEMASE LIKE PROTEIN	Epimerase
17144	2146	REVERSE TRANSCRIPTASE LIKE PROTEIN	Transcriptase

17154	2147	ACYLAMINOACYL-PEPTIDASE LIKE PROTEIN	Protease
17160	2148	GERMIN PRECURSOR OXALATE OXIDASE	Oxidase
17163	2149	RNA POLYMERASE II FIFTH LARGEST SUBUNIT LIKE	Polymerase
		PROTEIN	
17188	2150	IMIDAZOLEGLYCEROL-PHOSPHATE DEHYDRATASE	Dehydratase
17218	2151	ABC TRANSPORTER HOMOLOG	Transporter
17222	2152	GLUCOSYLTRANSFERASE	Transferases
17223	2153	UTP-GLUCOSE GLUCOSYLTRANSFERASE LIKE	Transferases
		PROTEIN	
17224	2154	UTP-GLUCOSE GLUCOSYLTRANSFERASE	Transferases
17225	2155	CELLULOSE SYNTHASE LIKE PROTEIN	Synthase
17228	2156	CELLULOSE SYNTHASE LIKE PROTEIN	Synthase
17240	2157	HYDROPEROXIDE LYASE (HPOL) LIKE PROTEIN	Lyase
17244	2158	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		LIKE PROTEIN .	
17245	2159	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		LIKE PROTEIN	
17246	2160	INDOLE-3-ACETATE BETA-GLUCOSYLTRANSFERASE	Transferases
		LIKE PROTEIN	
17249	2161	PYRUVATE PHOSPHATE DIKINASE,	Kinase
		PYRUVATE,ORTHOPHOSPHATE DIKINASE	
17251	2162	GLUCOSYLTRANSFERASE LIKE PROTEIN	Transferases
17261	2163	KINASE LIKE PROTEIN	Kinase, Protein
17283	2164	DELTA-CADINENE SYNTHASE LIKE PROTEIN	Synthase
17294	2165	PECTINESTERASE LIKE PROTEIN	Esterase
17309	2166	GALACTOKINASE LIKE PROTEIN	Kinase
17322	2167	BETA-1,3-GLUCANASE CLASS I PRECURSOR	Glycosylase
17323	2168	PEROXIDASE LIKE PROTEIN	Oxidase
17327	2169	NARINGENIN 3-DIOXYGENASE LIKE PROTEIN	Oxygenases
17353	2170	CELLULOSE SYNTHASE LIKE PROTEIN	Synthase
17354	2171	GLUCOSYLTRANSFERASE LIKE PROTEIN	Transferases
17363	2172	CYANOHYDRIN LYASE LIKE PROTEIN	Lyase
17367	2173	LIMONENE CYCLASE LIKE PROTEIN	Cyclase
17368	2174	LIMONENE CYCLASE LIKE PROTEIN	Cyclase
		Landa de la companya de la companya de la companya de la companya de la companya de la companya de la companya	

17371	2175	GIBBERELLIN OXIDASE-LIKE PROTEIN	Oxidase
17376		TRIACYLGLYCEROL LIPASE LIKE PROTEIN	Lipase
17390	2177	KINASE LIKE PROTEIN	Kinase, Protein
17402	2178	BETA-AMYLASE [PUTATIVE]	Glycosylase
17403	2179	SERINE PROTEASE-LIKE PROTEIN [PUTATIVE]	Protease
17427	2180	PHOSPHORIBOSYLGLYCINAMIDE	Transferases
		FORMYLTRANSFERASE-LIKE PROTEIN	
17464	2181	PEROXIDASE LIKE PROTEIN	Oxidase
17477	2182	N-ACETYLORNITHINE DEACETYLASE-LIKE PROTEIN,	
		FRAGMENT N-ACETYLORNITHINE DEACETYLASE	1
		(AODD) - DICTYOSTELIUM DISCOIDEUM, PID:G763048	
17510	2183	POTASSIUM CHANNEL - LIKE PROTEIN KCOI,	Channel
		ARABIDOPSIS THALIANA, Y07825	
17512	2184	POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
		EXOPOLYGALACTURONASE, ARABIDOPSIS	
		THALIANA, PIR2:S34266	
17518	2185	STARCH SYNTHASE-LIKE PROTEIN BACTERIAL AND	Synthase
		PLANT GLYCOGEN (STARCH) SYNTHASES; FOR	
		EXAMPLE B.SUBTILIS, PATCHX:D1020368	
17519	2186	RECEPTOR SERINE/THREONINE KINASE-LIKE PROTEIN	Kinase, Protein
		RECEPTOR SERINE/THREONINE KINASE PR5K,	
		PATCHX:G1235680	
17521	2187	PHOSPHO-N-ACETYLMURAMOYL-PENTAPEPTIDE-	Transferases
		TRANSFERASE, HAEMOPHILUS	
		INFLUENZAE,PIR2:A64185[PUTATIVE]	
17523	2188	POTASSIUM CHANNEL PROTEIN KAT2	Channel
17528	2189	BETA-1,3-GLUCANASE-LIKE PROTEIN STRONG	Glycosylase
		SIMLARITY TO ENDO-BETA-1,3-BETA-D-	
		GLUCOSIDASE, NICOTIANA TABACUM, PIR2:S46495	
17538	2190	ADENYLOSUCCINATE LYASE - LIKE PROTEIN	Lyase
		ADENYLOSUCCINATE LYASE - HAEMOPHILUS	
		INFLUENZAE	
17549	2191	LIPASE-LIKE PROTEIN PN47P, IPOMOEA NIL,	Lipase
		PID:G1527001	
17572	2192	CELLULOSE SYNTHASE - LIKE PROTEIN CELLULOSE	Synthase
		SYNTHASE, GOSSYPIUM HIRSUTUM, U58283	
<b></b>			

17576	2193	DNA POLYMERASE III HOLOENZYME TAU SUBUNIT.	Polymerase
		THERMUS THERMOPHILUS, GB:AF025391[PUTATIVE]	
17589	2194	PROTEIN KINASE - LIKE PROTEIN PROTEIN KINASE 6,	Kinase, Protein
		GLYCINE MAX., PIR2:S29851	
17593	2195	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE - LIKE	Glycosylase
		PROTEIN XYLOGLUCAN ENDO-TRANSGLYCOSYLASE	
		RELATED PROTEIN XTR-4, ARABIDOPSIS THALIANA,	
		PIR2:S71223	
17596	2196	DNA (CYTOSINE-5-)-METHYLTRANSFERASE,	Transferases
		ARABIDOPSIS THALIANA, PIR2:S59604[PUTATIVE]	
17635	2197	PECTINACETYLESTERASE PROTEIN	Esterase
		PECTINACETYLESTERASE PRECURSOR, VIGNA	
		RADIATA, PIR2:S68805[PUTATIVE]	
17636	2198	PECTINACETYLESTERASE PECTINACETYLESTERASE	Esterase
		PRECURSOR, VIGNA RADIATA, PIR2:S68805[PUTATIVE]	
17662	2199	FE(II) TRANSPORT PROTEIN FE(II) TRANSPORT	Transporter
		PROTEIN, ARABIDOPSIS THALIANA,	
15.50	2222	GB:U27590[PUTATIVE]	
17663		FE(II) TRANSPORT PROTEIN	Transporter
17665	2201	ASPARTATE KINASE-HOMOSERINE DEHYDROGENASE - LIKE PROTEIN AK-HSDH BIFUNCTIONAL ENZYME	Denydrogenases
		PRECURSOR, ARABIDOPSIS THALIANA, PIR2:S46497	
17666	2202	CHITINASE / LYSOZYME PZ PRECURSOR, NICOTIANA	Chitinggo
17000	2202	TABACUM, PIR2:S51591[PUTATIVE]	Cintinase
17668	2203	CHITINASE / LYSOZYME PZ PRECURSOR, NICOTIANA	Chitinase
1,000	2203	TABACUM, PIR2:S51591[PUTATIVE]	·
17677	2204		Isomerase
		TYPE, SYNECHOCYSTIS SP., PIR2:S75144[PUTATIVE]	
17688	2205	POTASSIUM TRANSPORTER-LIKE PROTEIN PUTATIVE	Transporter
		POTASSIUM TRANSPORTER ATKT2P & ATKT1P,	-
		ARABIDOPSIS THALIANA, PATCHX:G2384669 &	}
		PATCHX:G2384671	
		·	
17700	2206	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE	Transferases
17700	ļ	PHOSPHORIBOSYLANTHRANILATE TRANSFERASE, PHOSPHORIBOSYLANTHRANILATE TRANSFERASE,	Transferases
17700			Transferases

17703	2207	VACUOLAR SORTING RECEPTOR-LIKE PROTEIN BP-80 Receptor
1		VACUOLAR SORTING RECEPTOR, PISUM SATIVUM,
		PATCHX:G1737222
17712	2208	TERPENE CYCLASE LIKE PROTEIN 5-EPI-Cyclase
		ARISTOLOCHENE SYNTHASE, NICOTIANA TABACUM,
		PATX:G505588
17713	2209	CADINENE SYNTHASE LIKE PROTEIN (+)-DELTA-Synthase
		CADINENE SYNTHASE ISOZYME XC14,
		GOSSYPIUMARBOREUM, PIR2:S68366
17715	2210	TERPENE CYCLASE LIKE PROTEIN VETISPIRADIENE Cyclase
		SYNTHASE, HYOSCYAMUS MUTICUS, PATX:G763421
17719	2211	CLV1 RECEPTOR KINASE LIKE PROTEIN CLAVATAI Kinase, Protein
	····	RECEPTOR KINASE, ARABIDOPSIS TH., PATX:G2160756
17744	2212	RNA-DIRECTED DNA POLYMERASE, ARABIDOPSIS Polymerase
		THALIANA, PIR:S65815[PUTATIVE]
17745	2213	PROTEIN KINASE LIKE PROTEINS, Kinase, Protein
		ARABIDOPSIS[PUTATIVE]
17746	2214	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17747	2215	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17748	2216	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
15540		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17749	2217	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
15550		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17750		RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
15551		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17751		RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
17750		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17752	2220	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
17750	2221	PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17753		RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
1885		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]
17754		RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1, Kinase, Protein
		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]

17755	2223	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1,	Kinase, Protein
		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]	,
17756	2224	RECEPTOR-LIKE PROTEIN KINASE HOMOLOG RK20-1,	Kinase, Protein
		PHASEOLUS VULGARIS, GB:AF078082[PUTATIVE]	
17757	2225	PROTEIN (FRAGMENT) RECEPTOR-LIKE PROTEIN	Kinase, Protein
		KINASE HOMOLOG RK20-1, PHASEOLUS VULGARIS,	
		GB:AF078082[PUTATIVE]	
17758	2226	RECEPTOR-LIKE PROTEIN KINASE, RLK3,	Kinase, Protein
		ARABIDOPSIS THALIANA, AJ011674[PUTATIVE]	
17759	2227	PROTEIN KINASE LIKE PROTEINS,	Kinase, Protein
		ARABIDOPSIS[PUTATIVE]	
17761	2228	PROTEIN KINASE LIKE PROTEINS, ARABIDOPSIS	Kinase, Protein
		THALIANA[PUTATIVE]	
17762	2229	PROTEIN KINASE LIKE PROTEINS,	Kinase, Protein
		ARABIDOPSIS[PUTATIVE]	
17770	2230	RECEPTOR-LIKE PROTEIN KINASE, RLK3,	Kinase, Protein
ų.		ARABIDOPSIS THALIANA, AJ011674[PUTATIVE]	
17773	2231	RETICULINE OXIDASE - LIKE PROTEIN RETICULINE	Oxidase
		OXIDASE (EC 1.5.3.9) PRECURSOR, ESCHSCHOLZIA	
		CALIFORNICA, PIR2:A41533	
17774	2232	RETICULINE OXIDASE -LIKE PROTEIN RETICULINE	Oxidase
		OXIDASE, ESCHSCHOLZIA CALIFORNICA, PIR:A41533	:
17775	2233	RETICULINE OXIDASE - LIKE PROTEIN RETICULINE	Oxidase
		OXIDASE PRECURSOR, ESCHSCHOLZIA CALIFORNICA,	
		PIR:A41533	
17776	2234	TRIPEPTIDYL-PEPTIDASE II, HOMO SAPIENS, I	Protease
		PIR:S54376[PUTATIVE]	
17785	2235	RECEPTOR KINASE-LIKE GENE, ORYZA	Kinase, Protein
		LONGISTAMINATA, GB:U72725[PUTATIVE]	
17790	2236	CARBONIC ANHYDRASE -LIKE PROTEIN CARBONIC	Anhydrase
		ANHYDRASE, ARABIDOPSIS THALIANA, GB:U73462	
17791	2237	CARBONIC ANHYDRASE - LIKE PROTEIN DIOSCORIN	Anhydrase
		CLASS A PRECURSOR, DIOSCOREA CAYENENSIS,	
		PIR:S57766	
	1	DID. C57766	

17811	2238	GIBBERELLIN 20-OXIDASE - LIKE PROTEIN	Oxidase
		GIBBERELLIN C-20 OXIDASE, ORYZA SATIVA	,
		PATCHX:G1854637	
17813	2239	UDP-3-O-[3-HYDROXYMYRISTOYL] GLUCOSAMINE N	Transferases
		ACYLTRANSFERASE, E.COLI, PIR2:S13729[PUTATIVE]	ł
17814	2240	RECEPTOR KINASE - LIKE PROTEIN SRK3 RECEPTOR	Kinase, Protein
		KINASE, BRASSICA OLERACEA, PATCHX:G624943	
17827	2241	RECEPTOR KINASE - LIKE PROTEIN RECEPTOR-LIKE	Kinase, Protein
		KINASE SFR2, BRASSICA OLERACEA, PID:E258943	
17828	2242	RECEPTOR-LIKE SERINE/THREONINE PROTEIN KINASE	Kinase, Protein
		ARK3	
17829	2243	SERINE/THREONINE KINASE - LIKE PROTEIN	Kinase, Protein
		SERINE/THREONINE KINASE BRLK, BRASSICA	
		OLERACEA, GB:Y12531	
17830	2244	SERINE/THREONINE PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR PROTEIN KINASE (IRK1), IPOMOEA	·
		TRIFIDA, GB:U20948	
17831	2245	SERINE/THREONINE KINASE - LIKE PROTEIN	Kinase, Protein
17839	2246	NADH DEHYDROGENASE LIKE PROTEIN NADH	Dehydrogenase
		DEHYDROGENASE (UBIQUINONE) (EC 1.6.5.3)CHAIN	
		NDII YEAST,PIR2:S26704 AND OTHER NADH	
		DEHYDROGENASES	
17849	2247	BIFUNCTIONAL NUCLEASE, ZINNIA ELEGANS,	Nuclease
[ [		GB:U90266[PUTATIVE]	
17850	2248	BIFUNCTIONAL NUCLEASE D, ZINNIA ELEGANS,	Nuclease
		GB:U90266[PUTATIVE]	
17853	2249	SERINE PROTEASE - LIKE PROTEIN CUCUMISIN,	Protease
		CUCUMIS MELO, A55800	·
17854	2250	SUBTILISIN-LIKE PROTEASE SUBTILISIN-LIKE	Protease
		PROTEASE - LYCOPERSICON ESCULENTUM,AJ006378	
17858	2251	PEPTIDE TRANSPORTER - LIKE PROTEIN PEPTIDE	Transporter
		TRANSPORTER (PTR1) - HORDEUM VULGARE,AF023472	
17859	2252	GIBBERELLIN 3 BETA-HYDROXYLASE - LIKE PROTEIN	Hydroxylase
		3B-HYDROXYLASE, SOLANUM LYCOPERSICUM,	
		AB010992	
17886	2253	PEROXIDASE PRXR1	Oxidase

PCT/EP01/09892

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17891	2254	PECTINESTERASE LIKE PROTEIN PECTINESTERASE,	Esterase
		LYCOPERSICON ESCULENTUM, PATX:E312172	
17898	2255	PECTATE LYASE LIKE PROTEIN PECTATE LYASE,	Lyase
		ZINNIA ELEGANS, PATX:E283787	ļ
17899	2256	PECTATE LYASE LIKE PROTEIN PECTATE LYASE,	Lyase
		ZINNIA ELEGANS, PATX:E283787	}
17903	2257	SERINE/THREONINE PROTEIN KINASE LIKE PROTEIN	Kinase, Protein
		SERINE/THREONINE PROTEIN KINASE (ARSKI GENE),	
		ARABIDOPSIS THALIANA, PATCHX:G1066501	
17904	2258	RECEPTOR LIKE PROTEIN (FRAGMENT) ES43 PROTEIN,	Receptor
		BARLEY, PIR2:S44281	
17916	2259	ALTERNATIVE OXIDASE, MANGIFERA INDICA,	Oxidase
		PIR2:S45035[PUTATIVE]	
17949	2260	TREHALOSE-6-PHOSPHATE PHOSPHATASE - LIKE	Phosphatase
		PROTEIN TREHALOSE-6-PHOSPHATE PHOSPHATASE	
]		(ATTPPA), PID:G2944178	
17963	2261	LEUCINE RICH REPEAT RECEPTOR KINASE-LIKE	Kinase, Protein
		PROTEIN LEUCINE RICH REPEAT RECEPTOR-LIKE	
		KINASE, ORYZA SATIVA, PATCHX:E267533	
17974	2262	BILE ACID SODIUM-DEPENDENT TRANSPORTER,	Transporter
		HOMO SAPIENS, PIR2:138655, PREDICTED PROTEIN	
}		ILEAL	
17977	2263	ANTHOCYANIDIN SYNTHASE - LIKE PROTEIN	Oxygenases
		PUTATIVE LEUCOANTHOCYANIDIN DIOXYGENASE,	
		ARABIDOPSIS THALIANA, PID:G1575699	
17978	2264	LEUCOANTHOCYANIDIN DIOXYGENASE (LDOX)	Oxygenases
		[PUTATIVE]	
18000	2265	GAMMA-GLUTAMYLCYSTEINE SYNTHETASE	Synthase
18003	2266	PROTEIN KINASE - LIKE PROTEIN SERINE/THREONINE	Kinase, Protein
		KINASE - BRASSICA OLERACEA	
18004	2267	SERINE/THREONINE KINASE - LIKE PROTEIN KI	Kinase, Protein
		DOMAIN INTERACTING KINASE 1 (KIK1), ZEA MAYS	
18005	2268	SERINE/THREONINE KINASE - LIKE PROTEIN KI	Kinase, Protein
	•	DOMAIN INTERACTING KINASE 1 (KIK1), ZEA MAYS	
18008	2269	SERINE/THREONINE KINASE -LIKE PROTEIN	Kinase, Protein
		SERINE/THREONINE KINASE, BRASSICA OLERACEA	

SERINE/THREONINE KINASE, BRASSICA OLERACEA  18010 2271 SERINE /THREONINE KINASE - LIKE PROTEIN SERINE Kinase, Prote /THREONINE KINASE, BRASSICA OLERACEA  18011 2272 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18012 2273 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote TRIFIDA[PUTATIVE]
/THREONINE KINASE, BRASSICA OLERACEA  18011 2272 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18012 2273 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA
18011 2272 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18012 2273 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1: JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
SERINE/THREONINE KINASE, BRASSICA OLERACEA  18012 2273 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
18012 2273 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1: JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
SERINE/THREONINE KINASE, BRASSICA OLERACEA  18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
18013 2274 S-RECEPTOR KINASE 8 PRECURSOR, BRASSICA Kinase, Prote CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
CAMPESTRIS, PIR1:JC2481[PUTATIVE]  18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
18014 2275 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Prote SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
SERINE/THREONINE KINASE, BRASSICA OLERACEA  18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
18015 2276 PROTEIN KINASE - LIKE PROTEIN RECEPTOR KINASE Kinase, Prote 1, BRASSICA RAPA 18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
1, BRASSICA RAPA  18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
18016 2277 RECEPTOR PROTEIN KINASE, IPOMOEA Kinase, Prote
TRIFIDA[PUTATIVE]
18017 2278 SERINE/THREONINE KINASE SERINE/THREONINE Kinase, Protein
KINASE, BRASSICA OLERACEA
18018 2279 SERINE /THREONINE KINASE - LIKE PROTEIN SERINE Kinase, Protein
/THREONINE KINASE, BRASSICA OLERACEA
18019 2280 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Protein
SERINE/THREONINE KINASE, BRASSICA OLERACEAE
18020 2281 SERINE/THREONINE KINASE - LIKE PROTEIN KI Kinase, Protein
DOMAIN INTERACTING KINASE 1 (KIK1), ZEA MAYS
18021 2282 SERINE/THREONINE KINASE - LIKE PROTEIN KI Kinase, Protein
DOMAIN INTERACTING KINASE 1, ZEA MAYS
18022 2283 SERINE/THREONINE KINASE - LIKE PROTEIN Kinase, Protei
SERINE/THREONINE KINASE, BRASSICA OLERACEA
18024 2284 ACC OXIDASE, NICOTIANA GLUTINOSA[PUTATIVE] Oxidase
18054 2285 POTASSIUM TRANSPORT PROTEIN (TRH1) HIGH-Transporter
AFFINITY POTASSIUM TRANSPORT PROTEIN KUP1,
ARABIDOPSIS THALIANA, EMBL:AC004165[PUTATIVE]
18061 2286 V-ATPASE SUBUNIT G (VAG2 GENE) ATPase
18064 2287 RECEPTOR KINASE RECEPTOR-LIKE PROTEIN KINASE Kinase, Protei
- ARABIDOPSIS THALIANA RKL1,
PID:G4008006[PUTATIVE]

3.2.1.15) - AVOCADO, EMBL:X66426[PUTATIVE]  18089 2289 CELLULOSE SYNTHASE CATALYTIC SUBUNIT - LIKE SYNTHASE PROTEIN CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), ARABIDOPSIS THALIANA, GB:AF027173  18090 2290 CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]  18091 2291 CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), Synthase ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase ALDOLASE -ESCHERICHIA COLI,PID-G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE -Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POlymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter 18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]	18072	2288	POLYGALACTURONASE POLYGALACTURONASE (EC	CGlycosylase
PROTEIN CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), ARABIDOPSIS THALIANA, GB:AF027173  18090 2290 CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]  18091 2291 CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), Synthase ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase ALDOLASE -ESCHERICHIA COLI,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE -Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]				
PROTEIN CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), ARABIDOPSIS THALIANA, GB:AF027173  18090 2290 CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]  18091 2291 CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), Synthase ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase ALDOLASE -ESCHERICHIA COLI,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE -Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]	18089	2289		E Synthase
SUBUNIT (ATH-A), ARABIDOPSIS THALIANA, GB:AF027173  18090 2290 CELLULOSE SYNTHASE CATALYTIC SUBUNIT, Synthase ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]  18091 2291 CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase ALDOLASE -ESCHERICHIA COLL,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE -Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 IRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR LYASE (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter Transporter VULGARIS, PIR1:S51031[PUTATIVE]			į	1
GB:AF027173     18090   2290   CELLULOSE SYNTHASE   CATALYTIC   SUBUNIT, Synthase     ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]     18091   2291   CELLULOSE SYNTHASE   CATALYTIC   SUBUNIT (ATH-A), Synthase     ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]     18098   2292   2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC   ACID   Aldolase     ALDOLASE   -ESCHERICHIA     COLI,PID:G633197[PUTATIVE]     18104   2293   HYDROXYMUCONIC   SEMIALDEHYDE   HYDROLASE, Hydrolase     PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]     18128   2294   DIHYDROFOLATE   REDUCTASE   Reductase     SCHIZOSACCHAROMYCES   POMBE,PID:E1320950[PUTATIVE]     18155   2295   TRNA   ISOPENTENYLTRANSFERASE   TRNA   Transferases     ISOPENTENYLTRANSFERASE   (EC 2.5.1.8), YEAST     (SACCHAROMYCES   CEREVISIAE), PIR2:S67176[PUTATIVE]     18157   2296   ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR   Lyase     (CYSTEINE   SULPHOXIDE   LYASE), ONION, PIR2:S29301[PUTATIVE]     18168   2297   PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA,   Lyase     PATX:E209876[PUTATIVE]     18169   2298   DNA   POLYMERASE   III   LIKE   PROTEIN   DNA   Polymerase     POLYMERASE   III   GAMMA   SUBUNIT   AQUIFEX     AEOLICUS, PIR2:A70460     18171   2299   ABC-TYPE TRANSPORT PROTEINS[PUTATIVE]   Transporter     18178   2300   ACID   PHOSPHATASE   (EC 3.1.3.2)   PAP, PHASEOLUS   Phosphatase     VULGARIS, PIR1:S51031[PUTATIVE]   Transporter     18189   2301   ALPHA-AMYLASE   LIKE   PROTEIN   ALPHA-AMYLASE   GIycosylase				
ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]  18091 2291 CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), Synthase ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID ALDOLASE -ESCHERICHIA COLL,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE - Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POlymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:SS1031[PUTATIVE]				
18091   2291   CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A), Synthase   ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]     18098   2292   2,4-DiHYDROXYHEPT-2-ENE-1,7-DIOIC   ACID   Aldolase   ALDOLASE   -ESCHERICHIA   COLI,PID:G633197[PUTATIVE]     18104   2293   HYDROXYMUCONIC   SEMIALDEHYDE   HYDROLASE, Hydrolase   PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]     18128   2294   DIHYDROFOLATE   REDUCTASE   Reductase   SCHIZOSACCHAROMYCES   POMBE,PID:E1320950[PUTATIVE]     18155   2295   TRNA   ISOPENTENYLTRANSFERASE   TRNA   Transferases   ISOPENTENYLTRANSFERASE   (EC 2.5.1.8), YEAST (SACCHAROMYCES   CEREVISIAE), PIR2:S67176[PUTATIVE]     18157   2296   ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR   Lyase   (CYSTEINE   SULPHOXIDE   LYASE), ONION, PIR2:S29301[PUTATIVE]     18168   2297   PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase   PATX:E209876[PUTATIVE]     18169   2298   DNA   POLYMERASE   III   LIKE   PROTEIN   DNA   Polymerase   POLYMERASE   III   GAMMA   SUBUNIT   AQUIFEX   AEOLICUS, PIR2:A70460     18171   2299   ABC-TYPE TRANSPORT   PROTEINS[PUTATIVE]   Transporter   18178   2300   ACID   PHOSPHATASE (EC 3.1.3.2)   PAP, PHASEOLUS   Phosphatase   VULGARIS, PIR1:SS1031[PUTATIVE]	18090	2290	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]  18098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase			ARABIDOPSIS THALIANA, GB:AF027173[PUTATIVE]	
181098 2292 2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID Aldolase ALDOLASE -ESCHERICHIA COLI,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE - Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]	18091	2291	CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A)	Synthase
ALDOLASE COLI,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE REductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POlymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE]  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			ARABIDOPSIS THALIANA; GB:AF027173[PUTATIVE]	
COLI,PID:G633197[PUTATIVE]  18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, Hydrolase PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE REductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]	18098	2292	2,4-DIHYDROXYHEPT-2-ENE-1,7-DIOIC ACID	Aldolase
18104 2293 HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE, PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE - Reductase SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR LYASE (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]			ALDOLASE -ESCHERICHIA	
PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]  18128 2294 DIHYDROFOLATE REDUCTASE Reductase  SCHIZOSACCHAROMYCES  POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases  ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	]		COLI,PID:G633197[PUTATIVE]	
18128 2294 DIHYDROFOLATE REDUCTASE REductase  SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18104	2293	HYDROXYMUCONIC SEMIALDEHYDE HYDROLASE,	Hydrolase
SCHIZOSACCHAROMYCES POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			PSEUDOMONAS STUTZERI, AF039534[PUTATIVE]	
POMBE,PID:E1320950[PUTATIVE]  18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POlymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]	18128	2294	DIHYDROFOLATE REDUCTASE -	Reductase
18155 2295 TRNA ISOPENTENYLTRANSFERASE TRNA Transferases ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]			SCHIZOSACCHAROMYCES	
ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST (SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]			POMBE,PID:E1320950[PUTATIVE]	
(SACCHAROMYCES CEREVISIAE), PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter 18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18155	2295	TRNA ISOPENTENYLTRANSFERASE TRNA	Transferases
PIR2:S67176[PUTATIVE]  18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			ISOPENTENYLTRANSFERASE (EC 2.5.1.8), YEAST	
18157 2296 ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR Lyase (CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE]  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			(SACCHAROMYCES CEREVISIAE),	
(CYSTEINE SULPHOXIDE LYASE), ONION, PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			PIR2:S67176[PUTATIVE]	
PIR2:S29301[PUTATIVE]  18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18157	2296	ALLIIN LYASE ALLIIN LYASE (EC 4.4.1.4) PRECURSOR	Lyase
18168 2297 PECTATE LYASE PECTATE LYASE, MUSA ACUMINATA, Lyase PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter 18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase		,	(CYSTEINE SULPHOXIDE LYASE), ONION,	
PATX:E209876[PUTATIVE]  18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase		, 	PIR2:S29301[PUTATIVE]	
18169 2298 DNA POLYMERASE III LIKE PROTEIN DNA Polymerase POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter 18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18168	i	·	Lyase
POLYMERASE III GAMMA SUBUNIT - AQUIFEX AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			PATX:E209876[PUTATIVE]	
AEOLICUS, PIR2:A70460  18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase  VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18169			
18171 2299 ABC-TYPE TRANSPORT PROTEINS[PUTATIVE] Transporter  18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase  VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			POLYMERASE III GAMMA SUBUNIT - AQUIFEX	
18178 2300 ACID PHOSPHATASE (EC 3.1.3.2) PAP, PHASEOLUS Phosphatase VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase			AEOLICUS, PIR2:A70460	
VULGARIS, PIR1:S51031[PUTATIVE]  18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase		2299	ABC-TYPE TRANSPORT PROTEINS[PUTATIVE]	Transporter
18189 2301 ALPHA-AMYLASE - LIKE PROTEIN ALPHA-AMYLASE, Glycosylase	18178			Phosphatase
VIGNA MUNGO, PIR2:S10514	18189	1		Glycosylase
			VIGNA MUNGO, PIR2:S10514	

18197	2302	MAGNESIUM-PROTOPORPHYRIN IX	Transferases
10197	2302		1
		METHYLTRANSFERASE - LIKE PROTEIN MAGNESIUM-	ł
		PROTOPORPHYRIN IX METHYLTRANSFERASE,	
		SYNECHOCYSTIS SP, PIR2:S71781	
18199	2303	SUPEROXIDE DISMUTASE (EC 1.15.1.1)	Mutase
		(FE)(FRAGMENT)	
18204	2304	ACID PHOSPHATASE - LIKE PROTEIN ACID	Phosphatase
		PHOSPHATASE-1,LYCOPERSICON ESCULENTUM,	
		SWISS-PROT:P27061	
18214	2305	PECTINESTERASE GROUP I CITRUS SINENSIS,	Esterase
		PID:G2098705[PUTATIVE]	
18215	2306	PECTINESTERASE - CITRUS SINENSIS,	Esterase
		PID:G2098705[PUTATIVE]	
18219	2307	SRG1-LIKE PROTEIN STRONG HOMOLOGY TO SRG1	Oxidase
		PROTEIN, A NEW MEMBER OF THE FE(II)/ASCORBATE	
		OXIDASE SUPERFAMILY, PID:G479047	
18220	2308	SRG1-LIKE PROTEIN STRONG HOMOLOGY TO SRG1	Oxidase
		PROTEIN, A NEW MEMBER OF THE FE(II)/ASCORBATE	
		OXIDASE SUPERFAMILY, PID:G479047	
18226	2309	CLPC PROTEASE - SPINACIA	Protease
		OLERACEA,PID:G4105131[PUTATIVE]	
18228	2310	RECEPTOR KINASE-LIKE PROTEIN RECEPTOR-LIKE	Kinase, Protein
		PROTEIN KINASE, RLK3 - ARABIDOPSIS	
		THALIANA,PID:E1363211	
18231	2311	GIBBERELLIN 20-OXIDASE - ARABIDOPSIS THALIANA	Oxidase
18259	2312	BETA-CAROTENE HYDROXYLASE	Hydroxylase
18261	2313	GLUTAMINE CYCLOTRANSFERASE PRECURSOR - LIKE	Transferases
		PROTEIN GLUTAMINE CYCLOTRANSFERASE	
		PRECURSOR, CARICA PAPAYA, AF061240	•
18270		XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE (XTR-6)	Glycosylase
18271	2315	XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE	Glycosylase
		XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE (EC	-
		3.2.1) XTR-3 - ARABIDOPSIS THALIANA (FRAGMENT),	
		PIR2:S71222[PUTATIVE]	
		* *	

18284	2316	V-ATPASE G-SUBUNIT LIKE PROTEIN G SUBUNIT OF	ATPase
		  VACUOLAR-TYPE H+-ATPASE (VAGI) - ARABIDOPSIS	
		THALIANA, PID:E1330303	
18286	2317	PHOSPHATIDYLSERINE DECARBOXYLASE	Decarboxylase
		PHOSPHATIDYLSERINE DECARBOXYLASE (EC 4.1.1.65)	
]		2 - YEAST, PIR2:S64484[PUTATIVE]	
18287	2318	PEROXIDASE PEROXIDASE (EC 1.11.1.7) PRECURSOR,	Oxidase
		CATIONIC (CLONEPNC2) - PEANUT,	
		PIR2:B38265[PUTATIVE]	
18290	2319	PEROXIDASE PEROXIDASE ATP13A - ARABIDOPSIS	Oxidase
		THALIANA, PID:E264765[PUTATIVE]	
18316	2320	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE-	Kinase
		DEPENDENT PHOSPHOFRUCTO-1-KINASE	
		PYROPHOSPHATE-DEPENDENT PHOSPHOFRUCTO-1-	
		KINASE, PRUNUS ARMENIACA, U93272	
18317	2321	STEROID SULFOTRANSFERASE - LIKE PROTEIN	Transferases
		STEROID SULFOTRANSFERASE, BRASSICA NAPUS,	
		AF000307	
18322	2322	SUBTILISIN PROTEASE - LIKE SUBTILISIN-LIKE	Protease
		PROTEASE, ARABIDOPSIS THALIANA, AF055848	
18336	2323	CALCIUM-DEPENDENT PROTEIN KINASE, ORYZA	Kinase, Protein
		SATIVA, PIR2:S56652[PUTATIVE]	
18343	2324	RECEPTOR PROTEIN KINASE - LIKE PROTEIN CF-2.1,	Kinase, Protein
		SOLANUM PIMPINELLIFOLIUM	
18372	2325	BETA-1,3-GLUCANASE BETA-1,3-GLUCANASE	' '
		PRECURSOR - ORYZA SATIVA,PID:G4097948[PUTATIVE]	j
18379			Cyclase
18396			Synthase
18397	2328		Isomerase
		ASPERGILLUS NIGER,PID:G899149[PUTATIVE]	
18414		DIHYDROKAEMPFEROL 4-REDUCTASE, MEDICAGO	Reductase
		SATIVA, PIR2:S61416[PUTATIVE]	
18418	2330	RECEPTOR LIKE KINASE RECEPTOR-LIKE KINASE	Kinase, Protein
		SFR2, BRASSICA OLERACEA, PID:E258943[PUTATIVE]	
18419		RECEPTOR PROTEIN KINASE RECEPTOR PROTEIN	Kinase, Protein
		KINASE, IPOMOEA TRIFIDA,PID:G836954[PUTATIVE]	

18433	2332	PROTOCHLOROPHYLLIDE REDUCTASE PRECURSOR	Reductase
18438			
10430	2333	METHANOBACTERIUM	Transferases
10445	020.4	THERMOAUTOTROPHICUM, PID: G2621768 [PUTATIVE]	
18445	2334	UDP RHAMNOSE-ANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE - LIKE PROTEIN UDP	
		RHAMNOSEANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE, PETUNIA HYBRIDA	· · · · · · · · · · · · · · · · · · ·
18446	2335	UDP RHAMNOSEANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE - LIKE PROTEIN UDP	
		RHAMNOSEANTHOCYANIDIN-3-GLUCOSIDE	
		RHAMNOSYLTRANSFERASE, PETUNIA X HYBRIDA,	
		PIR2:S36655	
18449	2336	CARBOHYDRATE KINASE - LIKE PROTEIN	Kinase
		PHOSPHOFRUCTOKINASE, BABESIA CANIS, AJ223322	
18468	2337	PROTEIN PHOSPHATASE HOMOLOG (PPH1)	Phosphatase
18482	2338	PROTEIN KINASE LEPK7, LYCOPERSICON	Kinase, Protein
		ESCULENTUM, GB:U89684[PUTATIVE]	
18495	2339	ATPASE, HAEMATOBIA IRRITANS, U12392[PUTATIVE]	ATPase
18497	2340	PECTINESTERASE LIKE PROTEIN PECTINESTERASE,	Esterase
		LYCOPERSICON ESCULENTUM, Z94058	
18500	2341	DIACYLGLYCEROL KINASE ZETA, HOMO SAPIENS,	Kinase
		GB:U94905[PUTATIVE]	
18501	2342	DIACYLGLYCEROL KINASE (FRAGMENT) PUTATIVE	Kinase
		DIACYLGLYCEROL KINASE - ARABIDOPSIS THALIANA,	
		PID:G4454484[PUTATIVE]	
18510	2343	NADH DEHYDROGENASE NADH DEHYDROGENASE	Dehydrogenase
		(UBIQUINONE) (EC 1.6.5.3) CHAIN NDI1 - YEAST,	
		PIR2:S26704[PUTATIVE]	
18520	2344	(1-4)-BETA-MANNAN ENDOHYDROLASE MANNAN	Hydrolase
		ENDO-1,4-BETA-MANNOSIDASE (EC 3.2.1.78) -	
		TOMATO, PIR2:T04323[PUTATIVE]	
18523	2345	RECEPTOR PROTEIN KINASE LIKE PROTEIN LECTIN	Kinase, Protein
		RECEPTOR-LIKE SERINE/THREONINE KINASE LECRKI,	
		ARABIDOPSIS THALIANA, PIR2:S68589	

18528	2346	PROTEIN PHOSPHATASE 2C-LIKE PROTEIN PROTEIN	Phosphatase
		PHOSPHATASE 2C-FISSION YEAST, PIR2:S54297	
18537	2347	RECEPTOR-LIKE PROTEIN KINASE 5 PRECURSOR	Kinase, Protein
		(RLK5)	1
18544	2348	PROTEIN (FRAGMENT) PROTEIN KINASE XA21, ORYZA	Kinase, Protein
	1	SATIVA, PIR1:A57676[PUTATIVE]	<b>{</b>
18553	2349	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE 5, ARABIDOPSIS	
	<u> </u>	THALIANA, PIR1:S27756	
18555	2350	SERINE/THREONINE KINASE-LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE SERINE/THREONINE KINASE (RKF2),	
	İ	ARABIDOPSIS THALIANA, EMBL:AF024649	
18560	2351	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE-	Oxygenases
		FORMING), SUS SCROFA DOMESTICA,	
		PIR1:A33768[PUTATIVE]	
18573	2352	XYLOGLUCAN ENDOTRANSGLYCOSYLASE - LIKE	Glycosylase
		PROTEIN XYLOGLUCAN ENDOTRANSGLYCOSYLASE	
		1,FAGUS SYLVATICA, PID:E1354157	
18593	2353	SERINE/THREONINE-SPECIFIC KINASE LIKE PROTEIN	Kinase, Protein
		SERINE/THREONINE-SPECIFIC KINASE LECRK1 -	
		ARABIDOPSIS THALIANA, PIR2:S68589	
18597	2354	RNA-DIRECTED DNA POLYMERASE - ARABIDOPSIS	Polymerase
		THALIANA RETROTRANSPOSON TA11-1,	
		PIR2:S65812[PUTATIVE]	
18606	2355	SERINE/THREONINE-SPECIFIC RECEPTOR PROTEIN	Kinase, Protein
		KINASE SERINE/THREONINE-SPECIFIC RECEPTOR	
		PROTEIN KINASE - ARABIDOPSIS THALIANA,	
		PIR2:S71277[PUTATIVE]	
18610	2356	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE-	Kinase
		DEPENDENT PHOSPHOFRUCTO-1-KINASE-LIKE	
]		PROTEIN PYROPHOSPHATE-DEPENDENT	
		PHOSPHOFRUCTO-I-KINASE - PRUNUS ARMENIACA	
10614	2257	(APRICOT),PID:G2688822	Di i
18614			Phosphatase
		PHOSPHATASE-1 (EC 3.1.3) - LYCOPERSICON	
		ESCULENTUM,PIR2:T06587	

18615	2358	ACID PHOSPHATASE-LIKE PROTEIN ACIE	Phosphatase
		PHOSPHATASE-1 (EC 3.1.3) - LYCOPERSICON	1
		ESCULENTUM,PIR2:T06587	
18624	2359	BETA-1,3-GLUCANASE-LIKE PROTEIN BETA-1,3-	Glycosylase
		GLUCANASE (EC 3.2.1) 7 - GLYCINE MAX,PIR2:T05960	
18633	2360	SERINE/THREONINE-SPECIFIC RECEPTOR PROTEIN	Kinase, Protein
		KINASE-LIKE PROTEIN SERINE/THREONINE-SPECIFIC	-
		RECEPTOR PROTEIN KINASE LRRPK,PIR2:T08975	
18634	2361	PHOSPHOLIPASE A2-LIKE PROTEIN PUTATIVE	Lipase
		PHOSPHOLIPASE A2 - ORYZA SATIVA,PID:E1424908	
18635	2362	PHOSPHOLIPASE A2 - ORYZA	Lipase
		SATIVA,PID:E1424908[PUTATIVE]	
18642	2363	UDP-N-ACETYLGLUCOSAMINE O-ACYLTRANSFERASE	Transferases
		- LIKE PROTEIN UDP-N-ACETYLGLUCOSAMINE O-	
		ACYLTRANSFERASE, ALLOCHROMATIUM VINOSUM,	
		GB:L76417	
18645	2364	CYTIDINE DEAMINASE - LIKE PROTEIN CYTIDINE	Deaminase
		DEAMINASE 6, ARABIDOPSIS THALIANA, AF080676	
18646	2365	CYTIDINE DEAMINASE - LIKE PROTEIN CYTIDINE	Deaminase
		DEAMINASE 6, ARABIDOPSIS THALIANA, AF080676	
18648	2366	CYTIDINE DEAMINASE 7	Deaminase
18649	2367	CYTIDINE DEAMINASE 6 (CDA6)	Deaminase
18650	2368	CYTIDINE DEAMINASE 2 (CDA2)	Deaminase
18651	2369	CYTIDINE DEAMINASE 3 (CDA3)	Deaminase
18652	2370	CYTIDINE DEAMINASE 5 (CDA5)	Deaminase
18653	2371	CYTIDINE DEAMINASE 4 (CDA4)	Deaminase
18662	2372	CYTOKININ OXIDASE - LIKE PROTEIN CYTOKININ	Oxidase
		OXIDASE, ZEA MAYS, GB:Y18377	
18672	2373	THREONINE SYNTHASE	Synthase
18677	2374	CHOLINE MONOOXYGENASE - LIKE PROTEIN	Oxygenases
		CHOLINE MONOOXYGENASE PRECURSOR, SPINACIA	
		OLERACEA,GB:U85780	
18687	2375	SERINE/THREONINE-SPECIFIC RECEPTOR PROTEIN	Kinase, Protein
		KINASE LRRPK	
L			

18688	2376	DIHYDROPTEROATE SYNTHASE-LIKE PROTEIN	Synthase
		PROBABLE DIHYDROPTEROATE SYNTHASE - PISUM	
		SATIVUM,PIR2:T06595	
18705	2377		Oxidase
18715	_	XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE	
10/13	2310	1	Glycosylase
10516		PRECURSOR	
18716	2379	XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE-LIKE	Glycosylase
		PROTEIN XYLOGLUCAN ENDO-1,4-BETA-D-	
		GLUCANASE (EC 3.2.1) XTR-3 - ARABIDOPSIS	
		THALIANA,PIR2:S71222	
18717	2380	XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE-LIKE	Glycosylase
		PROTEIN XYLOGLUCAN ENDO-1,4-BETA-D-	:
		GLUCANASE (EC 3.2.1) XTR-3 - ARABIDOPSIS	
		THALIANA,PIR2:S71222	
18724	2381	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL-LIKE PROTEIN CYCLIC NUCLEOTIDE	
		AND CALMODULIN-REGULATED ION CHANNEL -	
		ARABIDOPSIS THALIANA,PID:E1421684	
18732	2382	NUCLEOTIDE SUGAR EPIMERASE-LIKE PROTEIN I	Epimerase
		NUCLEOTIDE SUGAR EPIMERASE - VIBRIO	
]		VULNIFICUS,PID:G3093975	
18735	2383	CINNAMOYL-COA REDUCTASE - LIKE PROTEIN	Reductase
]		CINNAMOYL-COA REDUCTASE, SACCHARUM	
		OFFICINARUM, GB:AJ231134	
18740	2384	RECEPTOR-LIKE KINASE HOMOLOG SOMATIC	Kinase, Protein
		EMBRYOGENESIS RECEPTOR-LIKE KINASE - DAUCUS	
		CAROTA,PID:G2224911	
18741	2385	ANILINE DIOXYGENASE (GMP SYNTHASE LIKE)	Oxygenases
		PROTEIN - ACINETOBACTER	33
		SP.,PID:D1013698[PUTATIVE]	1
18742		ANILINE DIOXYGENASE (GMP SYNTHASE LIKE)	)xvgenases
		PROTEIN - ACINETOBACTER	11, 501111303
		SP.,PID:D1013698[PUTATIVE]	
18743		ANILINE DIOXYGENASE (GMP SYNTHASE LIKE)	Dyuganasas
10/43	1	·	Aygenases
	1		
		SP.,PID:D1013698[PUTATIVE]	

18744	2388	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL-LIKE PROTEIN CYCLIC NUCLEOTIDE	
		AND CALMODULIN-REGULATED ION CHANNEL -	
		ARABIDOPSIS THALIANA,PID:E1421684	
18746	2389	2-ACYLGLYCEROPHOSPHOETHANOLAMINE	Transferases
		ACYLTRANSFERASE - AQUIFEX	
		AEOLICUS,PIR2:E70476[PUTATIVE]	
18783	2390	OMEGA-6 FATTY ACID DESATURASE (FAD6)	Desaturases
		CHLOROPLAST	
18798	2391	SERINE/THREONINE-SPECIFIC PROTEIN KINASE - LIKE	Kinase, Protein
		SERINE/THREONINE-SPECIFIC PROTEIN KINASE	
	 	PRO25, ARABIDOPSIS THALIANA, PIR2:A46373	
18799	2392	WALL-ASSOCIATED KINASE 1, ARABIDOPSIS	Kinase, Protein
		THALIANA, GB:AJ009696[PUTATIVE]	
18802	2393	1,3-BETA-GLUCANASE - LIKE PROTEIN 1,3-BETA-	Glycosylase
		GLUCANASE PRECURSOR, GOSSYPIUM HIRSUTUM,	
		PIR2:S72529	
18808	2394	DNA TOPOISOMERASE LIKE- PROTEIN BACILLUS	Isomerase
		SUBTILIS DNA TOPOISOMERASE I; PID:G520753	
18809	2395	PROTEIN KINASE-LIKE PROTEIN SERINE/THREONINE-	Kinase, Protein
		SPECIFIC PROTEIN KINASE, ARABIDOPSIS THALIANA,	
		PIR2:S38326	
18810	2396	RECEPTOR-LIKE KINASE, PETUNIA INFLATA,	Kinase, Protein
		PATCHX:G1931655[PUTATIVE]	
18812		RECEPTOR KINASE - LIKE PROTEIN RECEPTOR	Kinase, Protein
		KINASE, PETUNIA INFLATA, PATCHX:G498278	
18816		CATION TRANSPORT PROTEIN CHAC, ESCHERICHIA	Transporter
		COLI, PIR2:G64868, PREDICTED PROTEIN	
18846	]	CELLULOSE SYNTHASE, AGROBACTERIUM	Synthase
100.50		TUMEFACIENS, PIR2:139714[PUTATIVE]	
18858		Glutamate KAINATE RECEPTOR, RATTUS NORVEGICUS,	Receptor
100.55		PIR2:I53474[PUTATIVE]	
18863	1	PEROXIDASE - LIKE PROTEIN PEROXIDASE, SPINACIA	Oxidase
		OLERACEA, PID:G1781338	

18865	2402	MONOGALACTOSYLDIACYLGLYCEROL SYNTHASE - S-	ynthase
		LIKE PROTEIN MONOGALACTOSYLDIACYLGLYCEROL	
		SYNTHASE, CUCUMIS SATIVUS, PID:G1805254	
18878	2403	ANTHRANILATE N-T	ransferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,DIAN	
		THUS CARYOPHYLLUS, Z84384[PUTATIVE]	
18887	2404	SERINE/THREONINE PROTEIN KINASE LIKE PROTEINK	inase, Protein
		VARIOUS PREDICTED PROTEIN KINASES,	
		ARABIDOPSIS THALIANA	
18912	2405	MIXED-LINEAGE PROTEIN KINASE, HOMO SAPIENS, K	inase, Protein
		PIR:A53800[PUTATIVE]	
18917	2406	S-RECEPTOR KINASE -LIKE PROTEINK	inase, Protein
		SERINE/THREONINE-SPECIFIC PROTEIN KINASE PK10	
		PRECURSOR, ORYZA SATIVA, PIR2:S50767	
18918	2407	L-ASCORBATE PEROXIDASE - LIKE PROTEIN VARIOUS O	xidase
		L-ASCORBATE PEROXIDASES	
18923	2408	POLYGALACTURONASE (EC 3.2.1.15)PRECURSOR, GI	lycosylase
		LYCOPERSICON ESCULENTUM,	
		PIR2:A25534[PUTATIVE]	
18924	2409	EXOPOLYGALACTURONASE, ARABIDOPSIS GI	lycosylase
1000		THALIANA, PIR2:S34200[PUTATIVE]	د,
18927			nthase
18940	2411	i	xygenases
		PROTEIN DIMETHYLANILINE MONOOXYGENASE (N-	
		OXIDE-FORMING), SUS SCROFA DOMESTICA,	
10051	2412	PIR:A33768	
18951	2412	POTASSIUM CHANNEL PROTEIN ATKC POTASSIUM CH	nannel
10057	2412	CHANNEL - SOLANUM TUBEROSUM, PATX:E264595	
18957	2415	1	nase, Protein
19070	2414	ARABIDOPSIS THALIANA, PIR1:JQ1674[PUTATIVE]	
18970	Z414)	PYROPHOSPHATEFRUCTOSE-6-PHOSPHATE 1-Tr. PHOSPHOTRANSFERASE PYROPHOSPHATE	ansferases
		•	
		FRUCTOSE-6-PHOSPHATE 1-PHOSPHOTRANSFERASE	
		(EC 2.7.1.90) - ENTAMOEBA HISTOLYTICA,	
		PIR2:S68243[PUTATIVE]	

18981	2415	PROTEIN PHOSPHOPROTEIN PHOSPHATASE,	Phosphatasa
10,01	2713	ARABIDOPSIS THALIANA, PIR2:S55457[PUTATIVE]	Hospitatase
18988	2416	FE(II) TRANSPORT PROTEIN, ARABIDOPSIS THALIANA,	Transporter
10,000	2410	GB:U27590[PUTATIVE]	Transporter
10003	2417		
18993	2417	PYRUVATE DECARBOXYLASE-I (PDC1)	Decarboxylase
18997	2418	CYCLOPROPANE-FATTY-ACYL-PHOSPHOLIPID	Synthase
		SYNTHASE, ESCHERICHIA COLI,	
		PIR2:A44292[PUTATIVE]	
18998	2419	CYCLOPROPANE-FATTY-ACYL-PHOSPHOLIPID	Synthase
		SYNTHASE, ESCHERICHIA COLI,	
		PIR2:A44292[PUTATIVE]	
19008	2420	PECTINESTERASE - LIKE PROTEIN PECTINESTERASE,	Esterase
		PRUNUS PERSICA, X95991	
19009	2421	PECTINESTERASE - LIKE PROTEIN PECTIN	Esterase
		METHYLESTERASE-LIKE PROTEIN, ZEA MAYS, Y13285	
19019	2422	GLYCOGENIN GLUCOSYLTRANSFERASE (EC 2.4.1.186) -	Transferases
		HUMAN, PID:G1174167[PUTATIVE]	
19022	2423	DIHYDROKAEMPFEROL 4-REDUCTASE (EC 1.1.1.219) -	Reductase
		SYNECHOCYSTIS, PIR2:S75325[PUTATIVE]	
19028	2424	PEROXIDASE ATP17A -LIKE PROTEIN PEROXIDASE	Oxidase
		ATP17A -A.THALIANA,PID:E252638	
19029	2425	SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE -	Kinase, Protein
		LIKE PROTEIN SOMATIC EMBRYOGENESIS RECEPTOR-	
[		LIKE KINASE -DAUCUS CAROTA,PID:G2224911	
19030	2426	POLYGALACTURONASE(EC 3.2.1.15) PRECURSOR -	Glycosylase
		ERWINIA CAROTOVORA,PID:G42330[PUTATIVE]	
19032	2427	ABC-TYPE TRANSPORT PROTEIN SLL1623 -	Transporter
		SYNECHOCYSTIS,PIR2:S74812[PUTATIVE]	
19037	2428	2-DEHYDRO-3-DEOXYPHOSPHOHEPTONATE	Aldolase
		ALDOLASE	
19039	2429	POTASSIUM TRANSPORTER ATKT5P (ATKT5)	Transporter
		[PUTATIVE]	
19044	2430	CARBONATE DEHYDRATASE - LIKE PROTEIN	Dehydratase
		CARBONATE DEHYDRATASE PRECURSOR, SPINACIA	
		OLERACEA,PIR2:S28797	
<u> </u>		<del></del>	<del></del>

19054	2431	AMINOTRANSFERASE (ASPC FAMILY), AQUIFEX	Transferases
		AEOLICUS, PIR2:D70479[PUTATIVE]	
19063	2432	INOSITOL 1,3,4-TRISPHOSPHATE 5/6-KINASE,	Kinase
		ARABIDOPSIS THALIANA, PIR2:JC5401[PUTATIVE]	
19067	2433	BETA-XYLAN ENDOHYDROLASE -LIKE PROTEIN (1,4)-	Hydrolase
		BETA-XYLAN ENDOHYDROLASE ISOENZYME X-II,	
		HORDEUM VULGARE, GB:U59313	
19068	2434	(1,4)-BETA-XYLAN ENDOHYDROLASE, HORDEUM	Hydrolase
		VULGARE, PID:G1718238[PUTATIVE]	
19069	2435	XYLAN ENDOHYDROLASE ISOENZYME X-I, HORDEUM	Hydrolase
		VULGARE,PID:G1813595[PUTATIVE]	
19070	2436	XYLAN ENDOHYDROLASE ISOENZYME X-I, HORDEUM	Hydrolase
		VULGARE,PID:G1813595[PUTATIVE]	
19071	2437	(1,4)-BETA-XYLAN ENDOHYDROLASE, ISOENZYME X-I,	Hydrolase
		HORDEUM VULGARE, PID:G1718236[PUTATIVE]	
19072	2438	XYLAN ENDOHYDROLASE ISOENZYME X-I, HORDEUM	Hydrolase
		VULGARE,PID:G1813595[PUTATIVE]	
19073	2439	PEROXIDASE PEROXIDASE ATP12A, ARABIDOPSIS	Oxidase
		THALIANA, PID:E264763[PUTATIVE]	
19077	2440	PROLYL 4-HYDROXYLASE ALPHA(II)-SUBUNIT, MUS	Hydroxylase
		MUSCULUS, PIR2:I49135[PUTATIVE]	
19078	2441	PROTEIN PHOSPHATASE WIP1, HOMO SAPIENS,	Phosphatase
		PID:G2218063[PUTATIVE]	
19099	2442	GLUCOSYLTRANSFERASE -LIKE PROTEIN IMMEDIATE-	Transferases
		EARLY SALICYLATE-INDUCED	
		GLUCOSYLTRANSFERASE, NICOTIANA TABACUM,	
10100	0.140	PIR2:T03747	
19108	2443	RECEPTOR PROTEIN KINASE - LIKE PROTEIN PROTEIN	Kinase, Protein
10100	2444	KINASE TMKL1, ARABIDOPSIS THALIANA, PID:E353150	
19109	2444		Dehydrogenases
		PROTEIN CINNAMYL ALCOHOL DEHYDROGENASE,	
10104	2447	POPULUS DELTOIDES, PATCHX:G288753	Winner D
19124	2445	MYOSIN HEAVY CHAIN KINASE B, DICTYOSTELIUM	Kinase, Protein
]		DISCOIDEUM, U90946[PUTATIVE]	1

19130	2446	SERINE/THREONINE PROTEIN KINASE DAUCUS	Kinase, Protein
		CAROTA SOMATIC EMBRYOGENESIS RECEPTOR-LIKE	
		KINASE[PUTATIVE]	
19136	2447	SERINE/THREONINE PROTEIN KINASE DAUCUS	Kinase, Protein
		CAROTA SOMATIC EMBRYOGENESIS RECEPTOR-LIKE	
		KINASE, GB:U93048[PUTATIVE]	:
19137	2448	FATTY ACID ELONGASE 1, KETOACYL-COA	Synthase
		SYNTHASE, ARABIDOPSIS	
	<u> </u>	THALIANA,GB:U29142[PUTATIVE]	
19140	2449	ISOFLAVONE REDUCTASE - LIKE PROTEIN	Reductase
		ISOFLAVONE REDUCTASE HOMOLOG P3,	
		ARABIDOPSIS THALIANA, PIR2:S57613	
19157	2450	ARGININE DECARBOXYLASE SPE2	Decarboxylase
19171	2451	CHALCONE SYNTHASE - LIKE PROTEIN CHALCONE	Synthase
		SYNTHASE HOMOLOG PRCHS1, PINUS RADIATA,	
		GB:U90341	
19174	2452	AMIDASE - LIKE PROTEIN AMIDASE, BACILLUS	Amidase
		FIRMUS, GB:U64312	
19178	2453	PHOSPHOLIPASE C, LISTERIA MONOCYTOGENES,	Lipase
		PIR:A37204[PUTATIVE]	
19179	2454	PHOSPHATIDYL INOSITOL-SPECIFIC PHOSPHOLIPASE	Lipase
		C, LISTERIA SEELIGERI, GB:X97014[PUTATIVE]	
19186		ł	Oxidase
19187	2456	BETA-GALACTOSIDASE - LIKE PROTEIN BETA-D-	Glycosylase
		GALACTOSIDASE, MANGIFERA INDICA, PID:G2209358	
19189	2457	PROTEIN KINASE - LIKE LEUCINE RICH REPEAT	Kinase, Protein
		RECEPTOR-LIKE KINASE, ORYZA SATIVA,	
		PATCHX:E267533	
19201	2458	O-METHYLTRANSFERASE - LIKE PROTEIN CAFFEIC	Transferases
j		ACID O-METHYLTRANSFERASE, PINUS TAEDA,	
		GB:U39301	
19202	2459	O-METHYLTRANSFERASE - LIKE PROTEIN CAFFEIC	Transferases
		ACID O-METHYLTRANSFERASE, PINUS TAEDA,	
1000	0466	GB:U39301	<u> </u>
19204	2460	AMINO ACID PERMEASE - LIKE PROTEIN LILY MRNA,	I ransporter
		LILIUM LONGIFLORUM, GB:D21814	

19209	2461	PROTEIN KINASE APKI, ARABIDOPSIS THALIANA	, Kinase, Protein
		PIR2:S28615[PUTATIVE]	
19215	2462	LIGAND-GATED ION CHANNEL PROTEIN	Channel
		ARABIDOPSIS THALIANA (PREDICTED)[PUTATIVE]	
19216	2463	SUGAR TRANSPORTER PROTEIN SUGAR	Transporter
		TRANSPORTER, ARABIDOPSIS THALIANA	1
		DB_XREF=PID:G1495273[PUTATIVE]	
19228	2464	DIHYDROFLAVONOL-4-REDUCTASE, GERBERA X SP.,	Reductase
		PIR2:S35189[PUTATIVE]	
19231		THIOREDOXIN REDUCTASE (NADPH) 2	Reductase
19245	2466	PROTEIN KINASE - LIKE PROTEIN SERINE/THREONINE-	Kinase, Protein
		SPECIFIC PROTEIN KINASE APK1, ARABIDOPSIS	
		THALIANA, PIR2:S28615	
19249	2467	,	Transferases
:		ACETYLTRANSFERASE, ARABIDOPSIS THALIANA,	
		PIR2:S71207[PUTATIVE]	
19252	2468		Glycosylase
		LYCOPERSICON ESCULENTUM,	
		PID:G2459815[PUTATIVE]	
19264	2469	PHOSPHOLIPASE D, ARABIDOPSIS THALIANA,	Lipase
100.55	0.450	GB:U84568[PUTATIVE]	
19266	2470	PROLYL 4-HYDROXYLASE ALPHA (II) SUBUNIT, HOMO	Hydroxylase
100.55		SAPIENS, GB:U90441[PUTATIVE]	
19267	2471	PROCOLLAGEN-PROLINE DIOXYGENASE,	Oxygenases
10000	0.470	CAENORHABDITIS ELEGANS, PIR2:A55069[PUTATIVE]	
19282	2472	ASCORBATE PEROXIDASE ASCORBATE PEROXIDASE,	Oxidase
10000	0.470	GOSSYPIUM HIRSUTUM, GB:U37060[PUTATIVE]	
19302		RECEPTOR PROTEIN KINASE CF-2.1 LEUCINE RICH	Kinase, Protein
		REPEAT PROTEIN, SOLANUM PIMPINELLIFOLIUM, PATX:G1184075[PUTATIVE]	
10222		•	Ouidana
19322		PEROXIDASE LIKE PROTEIN	Oxidase
19356	24/5	GLUCOSYLTRANSFERASE-LIKE PROTEIN	Transferases
19360	2476	GERANYLGERANYL PYROPHOSPHATE SYNTHASE	Synthase
19368	2477	UDP-GLUCURONYLTRANSFERASE-LIKE PROTEIN	Transferases
19374	2478	MAP3K-LIKE PROTEIN KINASE	Kinase, Protein

19394	2479	HYDROXYNITRILE LYASE LIKE PROTEIN	Lyase
19395	2480	PECTINESTERASE LIKE PROTEIN	Esterase
19404	2481	RECEPTOR KINASE-LIKE PROTEIN	Kinase, Protein
19428	2482	RECEPTOR KINASE-LIKE PROTEIN	Kinase, Protein
19431	2483	BETA-KETOADIPATE ENOL-LACTONE HYDROLASE,	Hydrolase
		ACINETOBACTER SP., L05770[PUTATIVE]	
19438	2484	PEROXIDASE, PRXR2	Oxidase
19440	2485	PECTINESTERASE LIKE PROTEIN	Esterase
19442	2486	HYDROXYNITRILE LYASE LIKE PROTEIN	Lyase
19445	2487	FORMAMIDASE - LIKE PROTEIN FORMAMIDASE, METHYLOPHILUS METHYLOTROPHUS, PIR2: S74213	Amidase
19447	2488	FORMAMIDASE - LIKE PROTEIN FORMAMIDASE,	Amidase
		METHYLOPHILUS METHYLOTROPHUS, PIR2:S74213	
19452	2489	N-ACETYLTRANSFERASE HOOKLESS 1 [PROBABLE]	Transferases
19470	2490	AMINO-ACID N-ACETYLTRANSFERASE, ESCHERICHIA	Transferases
		COLI, PIR1:XYECAA[PUTATIVE]	
19475	2491	PHOSPHOENOLPYRUVATE CARBOXYKINASE (ATP) -	Kinase
		LIKE PROTEIN	
19478	2492	AMMONIUM TRANSPORTER SATI, GLYCINE MAX.,	Transporter
		AF069738[PUTATIVE]	
19482	2493	HEXOKINASE - LIKE PROTEIN HEXOKINASE,	Kinase
		ARABIDOPSIS THALIANA, PIR2:S71205	
19486	2494	ENDO-XYLOGLUCAN TRANSFERASE - LIKE PROTEIN	Transferases
		ENDO-XYLOGLUCAN TRANSFERASE, GOSSYPIUM	
		HIRSUTUM, D88413	
19490	2495	UDP-GLUCURONYLTRANSFERASE-LIKE PROTEIN	Transferases
19507	2496	CINNAMYL-ALCOHOL DEHYDROGENASE ELI3-1	Dehydrogenases
19508	2497	CINNAMYL-ALCOHOL DEHYDROGENASE ELI3-2	Dehydrogenases
19528	2498	CELLULOSE SYNTHASE - POPULUS ALBA X POPULUS	Synthase
		TREMULA (CEL1),PID:G3511285[PUTATIVE]	
19551	2499	PECTINESTERASE PECTINESTERASE - LYCOPERSICON	Esterase
		ESCULENTUM, PID:E312172[PUTATIVE]	

19554	2500	GERANYLGERANYL PYROPHOSPHATE SYNTHASE-	Synthase
		RELATED PROTEIN	
19560	2501	PROTEIN PHOSPHATASE-2C PROTEIN PHOSPHATASE-	Phosphatase
		2C (PP2C) - MESEMBRYANTHEMUM CRYSTALLINUM,	ļ
		PID:G3608412[PUTATIVE]	
19562	2502	MONOOXYGENASE 2 (MO2)	Oxygenases
19563	2503	PHOSPHOLIPASE LIKE PROTEIN ARABIDOPSIS	Lipase
		THALIANA PEARLI 4 MRNA, PID:G871782	
19564	2504	PHOSPHOLIPASE LIKE PROTEIN ARABIDOPSIS	Lipase
		THALIANA PEARLI 4 MRNA, PID:G871782	
19565	2505	PHOSPHATIDYLINOSITOL SYNTHASE	Synthase
		PHOSPHATIDYLINOSITOL SYNTHASE (PIS1) -	
		ARABIDOPSIS THALIANA, PID:E1313354[PUTATIVE]	
19567	2506	<u> </u>	Glycosylase
		EXOGALACTANASE (BETA-GALACTOSIDASE) -	
	· · · · · · · · · · · · · · · · · · ·	LYCOPERSICON ESCULENTUM,PID:E1363850	
19577	2507	PHOSPHOLIPASE C (EC 3.1.4.3)	Lipase
		PRECURSOR, PHOSPHATIDY LINOSITOL-SPECIFIC -	
10500	2500	LISTERIA MONOCYTOGENES, PIR2:A37204[PUTATIVE]	
19590	2508	RECEPTOR-LIKE PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE RLK3, ARABIDOPSIS	
19618	2500	THALIANA, AJ011674  RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE	Vince Dustein
19018	2309	PROTEIN KINASE, CATHARANTHUS ROSEUS,	Killase, Protein
	1	Z73295[PUTATIVE]	
19619	2510		Phosphatase
15015	2310	MONO-PHOSPHATASE, STREPTOMYCES ANULATUS,	Thosphalase
		X92429	
19628	2511	GLUCOSE-I-PHOSPHATE ADENYLYLTRANSFERASE	Transferases
		(APL3)	
19630	2512	NAD(P)H OXIDOREDUCTASE, ISOFLAVONE	Reductase
		REDUCTASE - LIKE PROTEIN PYRC2, PYRUS	
		COMMUNIS, AF071477	
19634	2513	RECEPTOR PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR PROTEIN KINASE ERECTA, ARABIDOPSIS	
		THALIANA	

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19639	2514	CINNAMYL-ALCOHOL DEHYDROGENASE CADI	Dehydrogenases
19641	2515	CELLULOSE SYNTHASE CATALYTIC SUBUNIT (ATH-A)	Synthase
19643	2516	UBIQUITIN SPECIFIC PROTEASE 66 - GALLUS	Protease
		GALLUS,PID:G3800764[PUTATIVE]	
19652	2517	MITOCHONDRIAL CARRIER - LIKE PROTEIN AGPET8,	Transporter
		ASHBYA GOSSYPII, EMBL:AJ006406	
19660	2518	SHIKIMATE KINASE - LIKE PROTEIN SHIKIMATE	Kinase
		KINASE PRECURSOR, LYCOPERSICON ESCULENTUM,	
		GB:S21584	
19704	2519	2-DEHYDRO-3-DEOXYPHOSPHOHEPTONATE	Aldolase
		ALDOLASE	
19722	2520	PROTEIN - KINASE PROTEIN KINASE TMK1 (EC 2.7.1),	Kinase, Protein
		RECEPTOR TYPE PRECURSOR, ARABIDOPSIS	
		THALIANA, PIR:JQ1674[PUTATIVE]	
19737	2521	ANTHRANILATE N-BENZOYLTRANSFERASE - LIKE	Transferases
		PROTEIN ANTHRANILATE N-BENZOYLTRANSFERASE,	
		CLOVE PINK, PIR:T10717	
19740	2522	LAX1 / AUX1 -LIKE PERMEASE	Transporter
19741	2523	ALPHA-1,4-N-ACETYLGLUCOSAMINYLTRANSFERASE,	Transferases
		HOMO SAPIENS, EMBL:AF141315[PUTATIVE]	
19742	2524	ALPHA AMYLASE [PUTATIVE]	Glycosylase
19743	2525	PHOSPHATASE (CTDP1), HOMO SAPIENS,	Phosphatase
		EMBL:AF154115; HYPOTHETICAL PROTEIN RNA	
		POLYMERASE II CTD	
19748	2526	PYRUVATE DECARBOXYLASE-LIKE PROTEIN	Decarboxylase
		PYRUVATE DECARBOXYLASE (EC 4.1.1.1) PDC1 -	
		ARABIDOPSIS THALIANA, PIR:T05315	
19749	2527	PYRUVATE DECARBOXYLASE-LIKE PROTEIN	Decarboxylase
		PYRUVATE DECARBOXYLASE (EC 4.1.1.1) PDC1 -	
	:	ARABIDOPSIS THALIANA, PIR:T05315	
19770	2528	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE - ARABIDOPSIS THALIANA,	
		EMBL:ATLECGENE	
19771	2529	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE - ARABIDOPSIS THALIANA,	
		EMBL:ATLECGENE	İ
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19772	2530	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE - ARABIDOPSIS THALIANA	ļ
		EMBL:ATLECGENE	
19774	2531	GAMMA-INTERFERON INDUCIBLE LYSOSOMAL THIOL	Reductase
1		REDUCTASE - HOMO	
		SAPIENS,EMBL:AF097362[PUTATIVE]	
19784	2532	TRANSPORTER PROTEIN NA+/H+ ANTIPORTER	Transporter
		PROTEINS[PUTATIVE]	
19785	2533	TRANSPORTER PROTEIN NA+/H+ ANTIPORTER	Transporter
		PROTEINS[PUTATIVE]	
19786	2534	PROTEIN PHOSPHATASE TYPE 2C - SACCHAROMYCES	Phosphatase
		CEREVISIAE, EMBL:U72346[PUTATIVE]	
19808	2535	PROTEIN KINASE - DIFFERENT SPECIES[PUTATIVE]	Kinase, Protein
19809	2536	(1-4)-BETA-MANNAN ENDOHYDROLASE-LIKE PROTEIN	Hydrolase
		MANNAN ENDO-1,4-BETA-MANNOSIDASE (EC 3.2.1.78) -	-
		LYCOPERSICON ESCULENTUM, PIR:T04323	
19811	2537	RECEPTOR PROTEIN KINASES[PUTATIVE]	Kinase, Protein
10922	2520	DECEMBER AND ACCOUNTS AND ACCOUNTS	7
19823	2538	PROTEIN KINASE WALL-ASSOCIATED KINASE 2 WAK2	Kinase, Protein
19833	2530	- ARABIDOPSIS THALIANA, EMBL:AJ012423[PUTATIVE] AMINO ACID TRANSPORT PROTEIN - ARABIDOPSIS	Transportor
19655	2339	THALIANA, EMBL:U39783[PUTATIVE]	Tansporter
19839	2540	HYDROLASE AT2G32150 - ARABIDOPSIS THALIANA,	Hydrolasa
15055	2240	EMBL:AC006223[PUTATIVE]	Trydiolase
19841	2541	RIBONUCLEASE II-LIKE PROTEIN RIBONUCLEASE II	Nuclease
		FAMILY PROTEIN, DEINOCOCCUS RADIODURANS,	r vaciouse
		PIR:C75571	
19843	2542	ABC TRANSPORTER -LIKE PROTEIN NBD-LIKE	Transporter
		PROTEIN POP, ARABIDOPSIS THALIANA,	Î
		EMBL:AF127664	ļ
19845	2543	SERINE/THREONINE-SPECIFIC PROTEIN KINASE NAK	Kinase, Protein
19846	2544	UBIQUITIN-PROTEIN LIGASE E3-ALPHA -LIKE PROTEIN	Ligase
	ľ	UBIQUITIN-PROTEIN LIGASE E3-ALPHA, MOUSE,	0
	i	PIR:T14318	
19856		PROTEIN PHOSPHATASE-2C, MESEMBRYANTHEMUM	Phosphatase
		CRYSTALLINUM, EMBL:AF079355[PUTATIVE]	F
	[		

19892	2546	PROTEIN PHOSPHATASE - LIKE PROTEIN PROTEIN	Phosphatase
	e.	PHOSPHATASE 2C HOMOLOG, MESEMBRYANTHEMUM	
		CRYSTALLINUM, EMBL:AF097667	
19896	2547	PROTEIN KINASE - LIKE PROTEIN KINASE APK2A,	Kinase, Protein
		ARABIDOPSIS THALIANA, EMBL:D88206	
19930	2548	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE	Kinase, Protein
		PROTEIN KINASE, ARABIDOPSIS THALIANA,	
		EMBL:ATLECGENE	
19943	2549	LYSINE DECARBOXYLASE - LIKE PROTEIN LYSINE	Decarboxylase
		DECARBOXYLASE, EIKENELLA CORRODENS,	
		EMBL:U89166	1_
19979	2550	MONODEHYDROASCORBATE REDUCTASE (NADH) -	
		LIKE PROTEIN MONODEHYDROASCORBATE	
		REDUCTASE (NADH), CUCUMBER, PIR:JU0182	
19986	2551	S-RECEPTOR KINASE - LIKE PROTEIN S-RECEPTOR	Kinase, Protein
		KINASE HOMOLOG PRECURSOR, RICE, PIR:S50767	
19990	2552	HISTONE DEACETYLASE -LIKE PROTEIN HISTONE	Esterase
		DEACETYLASE, HD2-P39, NUCLEOLAR, ZEA MAYS,	
10000		PIR:T04141	
19992	2553	BETA-(1-3)-GLUCOSYL TRANSFERASE,	Transferases
		BRADYRHIZOBIUM JAPONICUM,	
10003	2554	EMBL:AF047687[PUTATIVE] 3-DEOXY-D-MANNO-OCTULOSONIC ACID	T
19993	2554	TRANSFERASE -LIKE PROTEIN 3-DEOXY-D-MANNO-	Transferases
		OCTULOSONIC ACID TRANSFERASE, ESCHERICHIA	Ì
		COLI, PIR:JU0467	}
20014	2555	LIPASE -LIKE PROTEIN LIPASE ARAB-1, ARABIDOPSIS	Linase
2001.	2332	THALIANA, PIR:S68410	Dipuse
20028	2556		Mutase
20034		CARBONATE DEHYDRATASE - LIKE PROTEIN	Dehydratase
20047			Lyase
20053			Transferases
		CARBOXYL METHYLTRANSFERASE-LIKE PROTEIN	
20054	2560		Transferases
}		CARBOXYL METHYLTRANSFERASE-LIKE PROTEIN	
L			

20068	2561	2 OVOACVI LACVI CARRIER PROTERIE CVANTILACE	10 11
20008	2361		Synthase
		LIKE PROTEIN FABF 3-OXOACYL-[ACYL-CARRIER	1
		PROTEIN]SYNTHASE II, NEISSERIA MENINGITIDIS	,
		EMBL:U73942	
20069	2562	FATTY ACID ELONGASE - LIKE PROTEIN KCS1 FATTY	Synthase
		ACID ELONGASE 3-KETOACYL-COA SYNTHASE 1,	
		ARABIDOPSIS THALIANA, EMBL:AF053345	
20075	2563	SULPHITE REDUCTASE	Reductase
20110	2564	NICOTIANAMINE SYNTHASE (DBJ BAA74589.1)	Synthase
20111	2565	PECTINESTERASE	Esterase
20112	2566	PECTINESTERASE	Esterase
20132	2567	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
		(GB AAC39336.1)	
20142	2568	CHALCONE ISOMERASE, CHALCONE-FLAVONONE	Isomerase
		ISOMERASE [PUTATIVE]	
20147	2569	MONOOXYGENASE	Oxygenases
20149	2570	PEROXIDASE	Oxidase
20174	2571	PHOSPHORIBOSYLANTHRANILATE ISOMERASE	Isomerase
20175	2572	LEUCOANTHOCYANIDIN DIOXYGENASE-LIKE	Oxygenases
		PROTEIN	30
20178	2573	CATIONIC AMINO ACID TRANSPORTER 1[PUTATIVE]	Transporter
20188	2574	ANTHRANILATE SYNTHASE COMPONENT I-1	Synthase
		PRECURSOR (SP P32068)	
20201	2575	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN	Transferases
20202	2576	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN	Transferases
20203	2577	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN	Transferases
20204	2578	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN	Transferases
20205	2579	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN	Transferases
20231		SUCROSE TRANSPORTER PROTEIN	Transporter
20244	2581	LYSINE DECARBOXYLASE-LIKE PROTEIN	Decarboxylase
20248	2582	DIADENOSINE 5,5-P1,P4-TETRAPHOSPHATE	Hydrolase
1		HYDROLASE-LIKE PROTEIN	
20286		PEROXIDASE (EMB CAA68212.1)	Oxidase
		·	

20287	2584	PEROXIDASE	Oxidase
20288	2585	LECTIN-LIKE PROTEIN KINASE	Kinase, Protein
20289	2586	PROTEIN PHOSPHATASE 2C-LIKE	Phosphatase
20296	2587	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
20299	2588	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE- LIKE PROTEIN	Transferases
20300	2589	POLYGALACTURONASE INHIBITING PROTEIN 1; PGIP1 (GB AAF69827.1)	Glycosylase
20301	2590	POLYGALACTURONASE INHIBITING PROTEIN	Glycosylase
20307	2591	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
20313	2592	STEROID SULFOTRANSFERASE-LIKE PROTEIN	Transferases
20314	2593	STEROID SULFOTRANSFERASE-LIKE PROTEIN	Transferases
20321	2594	10-DEACETYLBACCATIN III-10-O-ACETYL TRANSFERASE - TAXUS CUSPIDATA, AF193765, EMBL:AF193765[PUTATIVE]	ĺ
20327	1	PROTEIN KINASE 6 - GLYCINE MAX, PIR:S29851[PUTATIVE]	Kinase, Protein
20328		PROTEIN KINASE XA21 - ORYZA SATIVA, PIR:A57676[PUTATIVE]	Kinase, Protein
20331		RECEPTOR-LIKE PROTEIN KINASE SEVERAL RECEPTOR-LIKE PROTEIN KINASES	Kinase, Protein
20333	2598	GIBBERELLIN 20-OXIDASE	Oxidase
20341		RECEPTOR-LIKE PROTEIN KINASE-LIKE PROTEIN RECEPTOR-LIKE PROTEIN KINASE - IPOMOEA NIL (JAPANESE MORNING GLORY), PIR:T18536	Kinase, Protein
20349	2600	AMIDASE [PUTATIVE]	AMIDASE
20350	- 1	INOSITOL HEXAKISPHOSPHATE KINASE 2 - HOMO SAPIENS, EMBL:AF177145[PUTATIVE]	Kinase
20354		PECTIN METHYL-ESTERASE-LIKE PROTEIN PECTIN METHYL-ESTERASE PER - MEDICAGO TRUNCATULA, EMBL:AJ249611	Esterase

20355	2603	PECTIN METHYL-ESTERASE-LIKE PROTEIN PECTIN	Esterase
		METHYL-ESTERASE PER - MEDICAGO TRUNCATULA	1
		EMBL:AJ249611	Ί
20356	2604	PECTIN METHYL-ESTERASE-LIKE PROTEIN PECTIN	Feterace
20330	2004	METHYL-ESTERASE PER - MEDICAGO TRUNCATULA.	
		EMBL:AJ249611	
20361	2605	2-OXOGLUTARATE-DEPENDENT DIOXYGENASE -	Oxygenases
20301	2003	SOLANUM CHACOENSE, EMBL:AF104925[PUTATIVE]	Oxygenases
20375	2606	RECEPTOR-LIKE PROTEIN KINASE[PUTATIVE]	Kinase, Protein
20373	2000	RECEFIOR-LIKE PROTEIN KINASE[FUTATIVE]	Kinase, Protein
20384	2607	EXONUCLEASE-LIKE PROTEIN	Nuclease
20385	2608	ALPHA GALACTOSYLTRANSFERASE PROTEIN	Transferases
20398	2609	PROANTHRANILATE N-BENZOYLTRANSFERASE -LIKE	Transferases
		PROTEIN ANTHRANILATE N-BENZOYLTRANSFERASE	
		(EC 2.3.1.144), DIANTHUS CARYOPHYLLUS,PIR:T10717	
20399	2610	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE -	Transferases
		LIKE PROTEIN N-	
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE,	
		IPOMOEA BATATAS, EMBL:AB035183	
20400	2611	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE -	Transferases
		LIKE PROTEIN ANTHRANILATE N-	
		BENZOYLTRANSFERASE, DIANTHUS CARYOPHYLLUS,	
		PIR:T10717	
20416	2612	GLYCEROPHOSPHODIESTER PHOSPHODIESTERASE -	Esterase
		LIKE PROTEIN GLYCEROPHOSPHODIESTER	
	!	PHOSPHODIESTERASE, BORRELIA HERMSII,	
		EMBL:BH40762	
20459	2613	LIPASE/HYDROLASE GDSL-like -motif	Lipase
20477	2614	FLAVONOL SYNTHASE (FLS) (SP Q96330)	Synthase
20480	2615	AMINO ACID TRANSPORT PROTEIN AAP2	Transporter
20496	2616	DNA-DIRECTED RNA POLYMERASE III CHAIN C53 -	Polymerase
		SACCHAROMYCES CEREVISIAE,	
		EMBL:X63501[PUTATIVE]	
20498	2617	POTASSIUM TRANSPORT PROTEIN-LIKE SEVERAL	Transporter
		POTASSIUM TRANSPORT PROTEINS	
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WO 02/10210 PCT/EP01/09892

20501	2610	HYDROLASE ETBD1 - RHODOCOCCUS SP.	TT-1-1-1-
20301	2018		Hydrolase
		EMBL:AB004320[PUTATIVE]	
20528	2619	BETA-GLUCOSIDASE - LIKE PROTEIN BETA-	Glycosylase
		GLUCOSIDASE, GLYCINE MAX, AF000378	
20531	2620	BETA-XYLOSIDASE - LIKE PROTEIN BETA-	Glycosylase
	<b> </b> 	XYLOSIDASE, HYPOCREA JECORINA, EMBL:Z69257	
20534	2621	PECTIN METHYLESTERASE - LIKE PROTEIN PECTIN	Esterase
		METHYLESTERASE, MELANDRIUM ALBUM,	
		EMBL:MAPME	
20545	2622	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
20560	2623	RECEPTOR PROTEIN KINASE -LIKE(FRAGMENT)	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE ERECTA,	
		ARABIDOPSIS THALIANA, EMBL:AC004484	
20563	2624	2,2-DIALKYLGLYCINE DECARBOXYLASE, P.CEPACIA,	Decarboxylase
Ì		EMBL:PCDGD[PUTATIVE]	
20568	2625	TREHALOSE-6-PHOSPHATE PHOSPHATASE -LIKE	Phosphatase
		PROTEIN TREHALOSE-6-PHOSPHATE PHOSPHATASE,	•
		ARABIDOPSIS THALIANA, EMBL:AF007779	
20574	2626	(3R)-HYDROXYMYRISTOYL-[ACYL CARRIER PROTEIN]	Dehydratase
		DEHYDRATASE -LIKE PROTEIN BETA-HYDROXYACYL-	-
		ACP DEHYDRATASE PRECURSOR, TOXOPLASMA	
		GONDII, EMBL:AF067150	
20577	2627		Transporter
ļ		ACINETOBACTER CALCOACETICUS,	r
		SWISSPROT:MUCK ACICA[PUTATIVE]	
20587		SERINE/THREONINE KINASE-LIKE PUTATIVE	Kinase Protein
	ł	SERINE/THREONINE KINASE - SORGHUM BICOLOR,	, 1 10tom
		EMBL:Y14600	ļ
20588			Tanaa
20308			Lyase
	ľ	ALPHA-HYDROXYNITRILE LYASE HNL4 - MANIHOT	
		ESCULENTA, EMBL:AJ223281	
20591		HISTIDINOL-PHOSPHATE AMINOTRANSFERASE-LIKE	Transferases
	j	PROTEIN HISTIDINOL-PHOSPHATE	
	ł	AMINOTRANSFERASE - NICOTIANA TABACUM,	
		EMBL:Y09204	

20606	2631	PROTEIN PHOSPHATASE PROTEIN TYROSINE	Phosphatase
		PHOSPHATASE-LIKE PROTEIN PTPLB, MUS	Ī
		MUSCULUS, EMBL:AF169286[PUTATIVE]	
20614	2632	BETA-XYLOSIDASE - LIKE PROTEIN BETA-	Glycosylase
		XYLOSIDASE, ASPERGLLUS NIDULANS,	
		EMBL:ANXLND	
20629	2633	HISTIDINE KINASE - LIKE PROTEIN SENSORY	Kinase, Protein
		TRANSDUCTION HISTIDINE KINASE SLR1759,	
		SYNECHOCYSTIS SP., PIR:S75142	1
20639	2634	LIGNOSTILBENE-ALPHA,BETA-DIOXYGENASE GENE,	Oxygenases
		SYNECHOCOCCUS PCC7942,	
		EMBL:AF055873[PUTATIVE]	
20644	2635	CHORISMATE MUTASE CM2	Mutase
20659	2636	SER/THR SPECIFIC PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
		PROTEIN KINASE 1, POPULUS NIGRA, EMBL:AB041503	
20668	2637	SUCROSE-PHOSPHATE SYNTHASE -LIKE PROTEIN	Synthase
		SUCROSE-PHOSPHATE SYNTHASE ISOFORM 1, CITRUS	
		UNSHIU, PIR:S72648	
20678	2638	GLUTAMATE RECEPTOR GLUR3 LIGAND-GATED	Channel
		CHANNEL-LIKE PROTEIN PRECURSOR, ARABIDOPSIS	
		THALIANA, EMBL:AF167355[PUTATIVE]	
20689	2639	FLAVIN-CONTAINING MONOOXYGENASE,	Oxygenases
		STREPTOMYCES COELICOLOR, PIR:T37052[PUTATIVE]	
20690	2640	SALICYLATE HYDROXYLASE, STREPTOMYCES	Hydroxylase
		COELICOLOR, PIR:T36193[PUTATIVE]	
20693	2641	SERINE/THREONINE PROTEIN KINASE, ARABIDOPSIS	Kinase, Protein
		THALIANA, SWISSPROT:NAK_ARATH[PUTATIVE]	
20695	2642	I-D-DEOXYXYLULOSE 5-PHOSPHATE SYNTHASE -	Synthase
		LIKE PROTEIN 1-D-DEOXYXYLULOSE 5-PHOSPHATE	
		SYNTHASE, LYCOPERSICON ESCULENTUM,	
		EMBL:AF143812	
20697	2643	SERINE THREONINE TYROSINE-SPECIFIC PROTEIN	Kinase, Protein
		KINASE APK1, ARABIDOPSIS THALIANA,	
		PIR:S28615[PUTATIVE]	
20711		L-GULONO-GAMMA-LACTONE OXIDASE, RATTUS	Oxidase
		NORVEGICUS, EMBL:RNFLAVIN[PUTATIVE]	

20714	2645	OLIGOPEPTIDE TRANSPORTER PROTEIN LEOPTI	Transporter
		OLIGOPEPTIDE TRANSPORTER, LYCOPERSICON	1 -
1		ESCULENTUM, EMBL:AF016713[PUTATIVE]	
20719	2646	MAP KINASE KINASE I, MUS MUSCULUS,	Kinase, Protein
		EMBL:AF117340[PUTATIVE]	
20722	2647	LYSOPHOSPHOLIPASE - LIKE PROTEIN	Lipase
		  LYSOPHOSPHOLIPASE HOMOLOG LPL1, ORYZA	•
		SATIVA, EMBL:AF039531	
20737	2648	POTASSIUM TRANSPORT PROTEIN GLUTATHIONE-	Transporter
		REGULATED POTASSIUM-EFFLUX SYSTEM PROTEIN	-
		KEFB, ESCHERICHIA COLI,	
		SWISSPROT:P45522[PUTATIVE]	
20745	2649	DIAMINOPIMELATE DECARBOXYLASE - LIKE PROTEIN	Decarboxylase
		DIAMINOPIMELATE DECARBOXYLASE, ARABIDOPSIS	
		THALIANA, EMBL:ATH249960	
20748	2650	ESTERASE - LIKE PROTEIN CINI PROTEIN - IMPORTED,	Esterase
		BUTYRIVIBRIO FIBRISOLVENS, PIR:T44624[PUTATIVE]	
20751	2651	SUBTILISIN-LIKE PROTEASE - LIKE PROTEIN	Protease
		SUBTILISIN-LIKE PROTEASE, ARABIDOPSIS	
		THALIANA, EMBL:AF098632	
20752	2652	LYSINE DECARBOXYLASE - LIKE PROTEIN LYSINE	Decarboxylase
		DECARBOXYLASE, EIKENELLA CORRODENS,	
		EMBL:U89166	
20766	2653	NPK1-RELATED PROTEIN KINASE[PUTATIVE]	Kinase, Protein
20770	2654	TRANSMEMBRANE TRANSPORT PROTEIN[PUTATIVE]	Transporter
20789	2655	GLUCOSYLTRANSFERASE -LIKE PROTEIN	Transferases
		GLUCOSYLTRANSFERASE IS5A, SALICYLATE-	
		INDUCED, COMMON TOBACCO, PIR:T03747	
20797	2656	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE -	Transferases
		LIKE PROTEIN ANTHRANILATE	
		PHOSPHORIBOSYLTRANSFERASE, GARDEN PEA,	
		PIR:T06460	
20800	2657	CALLOSE SYNTHASE CATALYTIC SUBUNIT -LIKE	Synthase
		PROTEIN PUTATIVE CALLOSE SYNTHASE CATALYTIC	
		SUBUNIT (CFL1), GOSSYPIUM HIRSUTUM,	}
		EMBL:AF085717	

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20805	2658	5-FORMYLTETRAHYDROFOLATE CYCLO-LIGASE-LIKE	1 -
		PROTEIN 5-FORMYLTETRAHYDROFOLATE CYCLO	-
		LIGASE (EC 6.3.3.2) - HOMO SAPIENS, EMBL:L38928	
20816	2659	PROTEIN KINASE-LIKE PROTEIN KINASE 1 PNPK1 -	Kinase, Protein
		POPULUS NIGRA, EMBL:AB041503	
20828	2660	ASPARTATE KINASE	Kinase
20829	2661	PROTEIN KINASE PRECURSOR - LIKE RECEPTOR-LIKE	Kinase, Protein
		PROTEIN KINASE 5 PRECURSOR, ARABIDOPSIS	
		THALIANA, SWISSPROT:RLK5_ARATH	
20841	2662	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE,	Isomerase
		SPODOPTERA FRUGIPERDA, EMBL:SF15038[PUTATIVE]	
20842	2663	TRANSALDOLASE - LIKE PROTEIN TRANSALDOLASE,	Aldolase
		SOLANUM TUBEROSUM, EMBL:U95923	
20870	2664	POLYAMINE OXIDASE	Oxidase
20871	2665	24-STEROL C-METHYLTRANSFERASE	Transferases
20874	2666	TRANSPORTER-LIKE PROTEIN	Transporter
20875	2667	TRANSPORTER-LIKE PROTEIN	Transporter
20887	2668	ENDOXYLOGLUCAN TRANSFERASE (DBJ BAA81669.1)	Transferases
20893	2669	CHALCONE SYNTHASE (NARINGENIN-CHALCONE	Synthase
		SYNTHASE) (TESTA 4 PROTEIN) (SP P13114)	
20898	2670	ALPHA-MANNOSIDASE	Glycosylase
20906	2671	ASPARTATE KINASE, LYSINE-SENSITIVE	Kinase
		(GB AAB63104.1)	
20910	2672	ABC TRANSPORTER, ATP-BINDING	Transporter
		PROTEIN[PUTATIVE]	
20913	2673	PEROXIDASE ATP20A (EMB CAA67338.1)	Oxidase
20921	2674	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
20922	2675	PROTOPORPHYRINOGEN IX OXIDASE	Oxidase
20926	2676	RIBULOSE-1,5-BISPHOSPHATE	Transferases
		CARBOXYLASE/OXYGENASE SMALL SUBUNIT N-	
		METHYLTRANSFERASE I, SPINACIA OLERACEA	
		CHLOROPLAST, PIR:T08996[PUTATIVE]	
20927	2677	KINASE - LIKE PROTEIN RING3 PROTEIN, HOMO	Kinase, Protein
		SAPIENS, EMBL:X96670	•

20931	2678	PROTEIN ESTERASE HDE, PETROLEUM-DEGRADING	Esterase
		BACTERIUM HD-1, EMBL:AB029896[PUTATIVE]	
20946	2679	TRNA SYNTHASE - LIKE PROTEIN TRNA	Synthase
		PSEUDOURIDINE SYNTHASE, FLAVOBACTERIUM	
		JOHNSONIAE, EMBL:AF169967	
20948	2680	N-ACETYLGLUCOSAMINYLTRANSFERASE III, MUS	Transferases
		MUSCULUS, EMBL:MMU66844[PUTATIVE]	
20957	2681	HIGH AFFINITY NITRATE TRANSPORTER - LIKE	Transporter
		PROTEIN HIGH AFFINITY NITRATE TRANSPORTER,	
		ORYZA SATIVA, EMBL:AB008519	
20962	2682	DNA METHYLTRANSFERASE 3, DANIO RERIO,	Transferases
		EMBL:AF135438[PUTATIVE]	
20965	2683	POLYGALACTURONASE - LIKE PROTEIN	Glycosylase
		POLYGALACTURONASE PG1, GLYCINE MAX,	
		EMBL:AF128266	
20970	2684	CINNAMOYL COA REDUCTASE - LIKE PROTEIN	Reductase
		CINNAMOYL COA REDUCTASE, POPULUS	
		TREMULOIDES, EMBL:AF217958	
20974			Anhydrase
20978	2686	FORMATE DEHYDROGENASE (FDH)	Dehydrogenases
20986	2687	GLUCOSYLTRANSFERASE -LIKE PROTEIN	ransferases
		GLUCOSYLTRANSFERASE IS5A, COMMON TOBACCO,	
		PIR:T03747	
20987	2688	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL -LIKE PROTEIN CYCLIC NUCLEOTIDE	
		AND CALMODULIN-REGULATED ION CHANNEL	
		CNGC6, ARABIDOPSIS THALIANA, EMBL:ATH17914	
20988	2689	CATION TRANSPORT PROTEIN PUTATIVE POTASSIUM	Fransporter
		TRANSPORTER ATKT2P, ARABIDOPSIS THALIANA,	
		EMBL:AF012657[PUTATIVE]	
20995	2690	ALPHA-MANNOSIDASE -LIKE PROTEIN MANNOSYL-	Glycosylase
		OLIGOSACCHARIDE 1,3-1,6-ALPHA-MANNOSIDASE,	
		MOUSE, PIR:A41641	
20998	2691	LYSOPHOSPHOLIPASE -LIKE PROTEIN	Lipase
		LYSOPHOSPHOLIPASE HOMOLOG (HU-K5), HUMAN,	
		EMBL:HSU67963	

21008	2692	SERINE/THREONINE SPECIFIC PROTEIN KINASE -LIKE	Kinase, Protein
		SERINE/THREONINE/TYROSINE-SPECIFIC PROTEIN	,
		KINASE APK1, ARABIDOPSIS THALIANA, PIR:S28615	
21009	2693	VOLTAGE-DEPENDENT ANION-SELECTIVE CHANNEL	Channel
		PROTEIN HSR2	}
21010	2694	AUXIN TRANSPORT PROTEIN - LIKE AUXIN	Transporter
		TRANSPORT PROTEIN (PIN7), ARABIDOPSIS	
		THALIANA, EMBL:AF087820	
21011	2695	PECTATE LYASE -LIKE PROTEIN PECTATE LYASE	Lyase
		LAT59, TOMATO, PIR:S27098	
21018	2696	PRX10 PEROXIDASE - LIKE PROTEIN PRX10	Oxidase
		PEROXIDASE, SPINACIA OLERACEA, EMBL:SOY16776	
21024	2697	AMINO ACID TRANSPORT PROTEIN, ARABIDOPSIS	Transporter
		THALIANA, EMBL:U39783[PUTATIVE]	
21038	2698	DNA METHYLTRANSFERASE 3, DANIO RERIO,	Transferases
		EMBL:AF135438[PUTATIVE]	
21039	2699	RRNA METHYLASE - LIKE PROTEIN RRNA METHYLASE	Methylase
		SPOU, AQUIFEX AEOLICUS, PIR:H70443	
21041	2700	CYCLIC NUCLEOTIDE-GATED CATION CHANNEL	Channel
21063	2701	PHYTOCHELATIN SYNTHETASE PUTATIVE	Synthase
		PHYTOCHELATIN SYNTHETASE - ARABIDOPSIS	
		THALIANA, EMBL:AJ006787[PUTATIVE]	
21064	2702	MITOCHONDRIAL CARRIER PROTEIN	Transporter
		MITOCHONDRIAL CARRIER PROTEIN - RIBES NIGRUM,	
		EMBL:AJ007580[PUTATIVE]	
21073	2703	SERINE/THREONINE-SPECIFIC PROTEIN KINASE-LIKE	Kinase, Protein
		PROTEIN SERINE/THREONINE-SPECIFIC PROTEIN	
		KINASE NPK15 - NICOTIANA TABACUM	· .
21086	2704	CARBOXYLESTERASE-LIKE PROTEIN STEROL	Esterase
		ESTERASE - RATTUS NORVEGICUS, EMBL:Z22803	
21087	ĺ	BETA-GLUCAN-ELICITOR RECEPTOR - GLYCINE MAX,	Receptor
		EMBL:D78510[PUTATIVE]	
21094	2706	PROTOCHLOROPHYLLIDE REDUCTASE HOMOLOG -	Reductase
		ORYZA SATIVA, EMBL:AF093628[PUTATIVE]	ļ

21100	2707	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
		PROBABLE SOMATIC EMBRYOGENESIS RECEPTOR	1
		LIKE KINASE - DAUCUS CAROTA, EMBL:U93048	
21101	2708	STEROID 5ALPHA-REDUCTASE-LIKE PROTEIN	Reductase
		STEROID 5ALPHA-REDUCTASE - RATTUS	S
		NORVEGICUS, PIR:A34239	
21112	2709	LIPASE-LIKE PROTEIN MONOGLYCERIDE LIPASE	Lipase
		MUS MUSCULUS, EMBL:AJ001118	
21119	2710	CELLULOSE SYNTHASE CELA - RHIZOBIUM	Synthase
		LEGUMINOSARUM, EMBL:AF121340[PUTATIVE]	
21123	2711	STEAROYL-ACYL CARRIER PROTEIN DESATURASE	Desaturases
1		STEAROYL-ACYL CARRIER PROTEIN DESATURASE	
		SAD1 - LINUM USITATISSIMUM, EMBL:AJ006957	}
21124	2712	STEAROYL-ACYL CARRIER PROTEIN DESATURASE	Desaturases
		STEAROYL-ACYL CARRIER PROTEIN DESATURASE	
		SAD1 - LINUM USITATISSIMUM, EMBL:AJ006957	
21129		ACETOLACTATE SYNTHASE-LIKE PROTEIN	Synthase
21139	2714	ACETYL-COA CARBOXYLASE, BIOTIN CARBOXYL	Carboxylase
		CARRIER PROTEIN OF ACETYL-COA CARBOXYLASE	
		PRECURSOR (BCCP) (SP Q42533)	
21141	2715		Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE[PUTA	
		TIVE]	
21145		S-ADENOSYLMETHIONINE:2-	Transferases
		DEMETHYLMENAQUINONE METHYLTRANSFERASE-	
		LIKE PROTEIN	
21148	2717	TYROSINE PHOSPHATASE[PUTATIVE]	Phosphatase
21150	2718	PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
21153	2719	AUXIN TRANSPORT PROTEIN[PUTATIVE]	Transporter
21159		RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
21169	1	CELLULASE (EC 3.2.1.4) PRECURSOR - XANTHOMONAS	Cellulase
		CAMPESTRIS PV. CAMPESTRIS, PIR:JH0158[PUTATIVE]	
21179	2722	N-ACETLYTRANSFERASE F13E7.7 - ARABIDOPSIS	Transferases
		THALIANA, EMBL:AC018363[PUTATIVE]	

21189	2723	RECEPTOR PROTEIN KINASE -LIKE PROTEIN	Kinase, Protein
		  SERINE/THREONINE-SPECIFIC RECEPTOR PROTEIN	1
	÷	KINASE (EC 2.7.1), ARABIDOPSIS THALIANA	,
		PIR:S71277	
21190	2724	CELLULOSE SYNTHASE CATALYTIC SUBUNIT -LIKE	Synthase
		PROTEIN ATH-B, CELLULOSE SYNTHASE CATALYTIC	:
		SUBUNIT, ARABIDOPSIS THALIANA, EMBL:AF027174	
21200	2725	SUGAR TRANSPORTER - LIKE PROTEIN D-XYLOSE-	Transporter
		PROTON SYMPORTER (D-XYLOSE TRANSPORTER)	
		LACTOBACILLUS BREVIS, SWISSPROT:XYLT_LACBR	
21202	2726	UDP GLUCOSE:FLAVONOID 3-O-	Transferases
		GLUCOSYLTRANSFERASE -LIKE PROTEIN UDP	
		GLUCOSE:FLAVONOID 3-O-GLUCOSYLTRANSFERASE,	
		VITIS VINIFERA, EMBL:AF000372	
21203	2727	· ·	Transferases
		GLUCOSYLTRANSFERASE -LIKE PROTEIN UDP	1 1
		GLUCOSE:FLAVONOID 3-O-GLUCOSYLTRANSFERASE,	
		VITIS VINIFERA, EMBL:AF000371	
21204	2728		Transferases
		GLUCOSYLTRANSFERASE -LIKE PROTEIN UDP	
		GLUCOSE:FLAVONOID 3-O-GLUCOSYLTRANSFERASE,	
21210	2720	VITIS VINIFERA, EMBL:AF000372	
21219		POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
21221	2/30	GLUTATHIONE S-TRANSFERASE-LIKE PROTEIN	Transferases
21222	2731	PHYTOENE SYNTHASE (GB AAB65697.1)	Synthase
21232	2732	GLUTAMATE DECARBOXYLASE I (GAD 1) (SP Q42521)	Decarboxylase
21241	2733	CELLULOSE SYNTHASE CATALYTIC SUBUNIT (IRX3)	Synthase
21252		PHOSPHOGLUCOMUTASE-LIKE PROTEIN	
21232		PHOSPHOGLUCOMUTASE, CHLOROPLAST - SPINACIA	Wittasc
		OLERACEA, EMBL:X75898	
21281	2735	PEROXIDASE ATP13A	Oxidase
21291		5-METHYLTETRAHYDROPTEROYLTRIGLUTAMATE	Transferases
		HOMOCYSTEINE S-METHYLTRANSFERASE	
21297		PHOSPHORIBOSYLANTHRANILATE TRANSFERASE-	Transferases
		LIKE PROTEIN	
	1		ŀ

21298	2738	ANTHRANILATE PHOSPHORIBOSYLTRANSFERASE	Transferases
		CHLOROPLAST PRECURSOR (SP Q02166)	
21311	2739	PROTEIN DISULFIDE ISOMERASE-RELATED	Isomerase
	  -	PROTEIN[PUTATIVE]	
21319	2740	GALACTOSE-1-PHOSPHATE URIDYL TRANSFERASE-	Transferases
,		LIKE PROTEIN	
21321	2741	BETA-1,3-GLUCANASE-LIKE PROTEIN	Glycosylase
21346	2742	RECEPTOR KINASE PK3 PRECURSOR, MAIZE,	Kinase, Protein
		PIR:T02753[PUTATIVE]	
21349	2743	SER/THR SPECIFIC PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
[ [		VARIOUS PROTEIN KINASE, ARABIDOPSIS THALIANA	
21360	2744	PROTEIN KINASE -LIKE PROTEIN PROTEIN KINASE 1,	Kinase, Protein
		POPULUS NIGRA, EMBL:AB041503	
21365	2745	PROTEIN 2'-HYDROXYISOFLAVONE REDUCTASE (EC	Reductase
		1.3.1.45) - NICOTIANA TABACUM,	
		PIR:T02202[PUTATIVE]	
21366	2746	BETA-AMYLASE-LIKE PROTEN BETA-AMYLASE -	Glycosylase
		PRUNUS ARMENIACA, EMBL:AF139501	
21390	2747	PROTEIN KINASE - LIKE PROTEIN PTO KINASE	Kinase, Protein
		INTERACTOR 1 (PTI1), LYCOPERSICON ESCULENTUM,	
		EMBL:SL28007	
21398	2748	PECTIN METHYL ESTERASE -LIKE PROTEIN PECTIN	Esterase
		METHYL ESTERASE, SOLANUM TUBEROSUM,	
		EMBL:AF152172	
21403	2749	TRNA ISOPENTENYLTRANSFERASE -LIKE PROTEIN	Transferases
		TRNA ISOPENTENYLTRANSFERASE,	
		SACCHAROMYCES CEREVISIAE, PIR:S67176	
21421	2750	GLUCOSE-1-PHOSPHATE ADENYLYLTRANSFERASE	Transferases
		(APL1/ADG2)	
21427	2751	PROTEIN PHOSPHATASE, KINASE ASSOCIATED	Phosphatase
21428	2752	PHOSPHOLIPASE - LIKE PROTEIN VARIOUS	Lipase
		PREDICTED PHOSPHOLIPASE PROTEINS	
21436	2753	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE - LIKE	Isomerase
	[:	PROTEIN PEPTIDYL-PROLYL CIS-TRANS ISOMERASE	
	.	A.THALIANA	
L	L		

21449	2754	PERMEASE FAMILY PROTEIN TC0205, CHLAMYDIA	Transporter
21119	2/54	MURIDARUM, PIR:D81729[PUTATIVE]	Tansporter
21457	2755	GLYOXAL OXIDASE PRECURSOR, PHANEROCHAETE	Ovidase
21437	2133	CHRYSOSPORIUM, PIR:A48296[PUTATIVE]	Oxidase
21462	2756	i	
21463	2756	PEPTIDE TRANSPORT PROTEIN-LIKE PEPTIDE	1 -
		TRANSPORT PROTEIN - HORDEUM VULGARE	·
		EMBL:AF023472	
21471	2757	GLUTAMINETRNA LIGASE - LUPINUS LUTEUS	Ligase
		EMBL:X91787[PUTATIVE, PROTEIN C-TERMINUS OF]	
21472	2758	PECTIN METHYLESTERASE-LIKE PROTEIN PECTIN	Esterase
		METHYL ESTERASE - SOLANUM TUBEROSUM,	
		EMBL:AF152172	
21487	2759	PEROXIDASE PEROXIDASE, LYCOPERSICON	Oxidase
		ESCULENTUM, PIR:S32768	
21488	2760	PEROXIDASE ATP N	Oxidase
21503	2761	IPP TRANSFERASE - LIKE PROTEIN TRNA DELTA(2)-	Transferases
		ISOPENTENYLPYROPHOSPHATE TRANSFERASE (IPP	
		TRANSFERASE), PSEUDOMONAS PUTIDA,	
		EMBL:AF016312	
21504	2762	RECEPTOR-LIKE PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE, ARABIDOPSIS	
		THALIANA, EMBL:M84659	
21527	2763	SUCROSE-PHOSPHATE SYNTHASE-LIKE PROTEIN	Synthase
		SUCROSE-PHOSPHATE SYNTHASE (EC 2.4.1.14)	
		ISOFORM 1 - CITRUS UNSHIU, EMBL:AB005023	
21532	2764	BETA-1,3-GLUCANASE BG4	Glycosylase
21533	2765	BETA-1,3-GLUCANASE BG5	Glycosylase
21538	2766	BETA-1,3-GLUCANASE-LIKE PROTEIN BETA-1,3-	Glycosylase
		GLUCANASE BG4 - A.THALIANA, EMBL:X79694	
21539	2767	ETHYLENE-FORMING-ENZYME-LIKE DIOXYGENASE-	Oxygenases
		LIKE PROTEIN ETHYLENE-FORMING-ENZYME-LIKE	
		DIOXYGENASE - PRUNUS ARMENIACA, EMBL:U97530	
21540	2768	MONOGALACTOSYLDIACYLGLYCEROL SYNTHASE	Synthase
21547	2769	RECEPTOR PROTEIN KINASE - LIKE PROTEIN KINASE	Kinase, Protein
		XA21, ORYZA SATIVA, PIR:A57676	

21554	2770	ETHYLENE-FORMING-ENZYME-LIKE DIOXYGENASE	- Oxygenases
		LIKE ETHYLENE-FORMING-ENZYME-LIKE	
		DIOXYGENASE, PRUNUS ARMENIACA, EMBL:U97530	j
21555	2771	BETA-1,3-GLUCANASE - LIKE PROTEIN BETA-1,3-	Glycosylase
		GLUCANASE BG4 AND BG5, A.THALIANA	,
		EMBL:ATCBG45	
21568	2772	RECEPTOR PROTEIN KINASE - LIKE RECEPTOR-LIKE	Kinase, Protein
}		PROTEIN KINASE PRK1, TOMATO, PIR:T07865	
21582	2773	SUCROSE-UDP GLUCOSYLTRANSFERASE	Transferases
21585	2774	PECTINESTERASE - LIKE PROTEIN PROBABLE	Esterase
		PECTINESTERASE PRECURSOR, GARDEN PEA,	[
		PIR:T06374	
21586	2775	BETA-1,3-GLUCANASE - LIKE PROTEIN PROBABLE	Glycosylase
		BETA-1,3-GLUCANASE, WHEAT, PIR:T06268	
21593	2776		Glycosylase
	······································	GLUCOSIDASE, COMMON NASTURTIUM, PIR:T10521	
21594	2777	BETA-D-GLUCAN EXOHYDROLASE - LIKE PROTEIN	,
		BETA-D-GLUCAN EXOHYDROLASE, NICOTIANA	
		TABACUM, EMBL:AB017502	
21597	2778		Transferases
		HOMOCYSTEINE S-METHYLTRANSFERASE - LIKE	
		PROTEIN 5-	
		METHYLTETRAHYDROPTEROYLTRIGLUTAMATE- HOMOCYSTEINE S-METHYLTRANSFERASE.	
		HOMOCYSTEINE S-METHYLTRANSFERASE, ARABIDOPSIS THALIANA, EMBL:U97200	
21605	2779	HOMOSERINE DEHYDROGENASE-LIKE PROTEIN THRA	Dehydrogenases
21005		BIFUNCTIONAL ENZYME - ESCHERICHIA COLI,	Denyarogenases
		PIR:B64720	
21634		PEROXIDASE ATP14A HOMOLOG	Oxidase
21635		ACYL COA REDUCTASE-LIKE PROTEIN	Reductase
21656	2782	PREPHENATE DEHYDRATASE/CHORISMATE MUTASE-	Dehydratase
		LIKE PROTEIN	
21658	2783	HISTONE DEACETYLASE-LIKE PROTEIN	Dehydratase
21667	2784	GLUCOSYLTRANSFERASE-LIKE PROTEIN	Transferases
21673	2785	LIPASE/HYDROLASE GDSL-like -motif	Linese
210/3	2103	EII ASE/II I DROLASE ODSL-IIKE -IIIOIII	Lipase

21677	2786	PROTEASE-LIKE PROTEIN	Protease
21682	2787	NA+/H+ ANTIPORTER-LIKE PROTEIN	Transporter
21683	2788	NA+/H+ ANTIPORTER-LIKE PROTEIN	Transporter
21688	2789	SERINE CARBOXYPEPTIDASE[PUTATIVE]	Protease
21693	2790	2-ISOPROPYLMALATE SYNTHASE-LIKE;	Synthase
		HOMOCITRATE SYNTHASE-LIKE	
21694	2791	2-ISOPROPYLMALATE SYNTHASE-LIKE PROTEIN	Synthase
21709	2792	SERINE/THREONINE PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
21736	2793	FERREDOXIN-THIOREDOXIN REDUCTASE VARIABLE	Reductase
		CHAIN[PUTATIVE]	
21751	2794	PHOSPHOTRANSFERASE[PUTATIVE]	Transferases
21759	2795	POLYA POLYMERASE[PUTATIVE]	Polymerase
21769	2796	GALACTINOL SYNTHASE	Synthase
21771	2797	AMINO ACID TRANSPORTER	Transporter
21776	2798	PECTINACETYLESTERASE	Esterase
21783	2799	ACYLTRANSFERASE	Transferases
21785	2800	DELTA-(+)CADINENE SYNTHASE (D-CADINENE	Synthase
		SYNTHASE)	
21790	2801	RECEPTOR-PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
21796	2802	PEROXIDASE-LIKE PROTEIN	Oxidase
21797	2803	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
21798	2804	ENDOCHITINASE ACIDIC (DBJ BAA21861.1)	Chitinase
21799	2805	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
21819	2806	STARCH SYNTHASE SOLUBLE	Synthase
21842	2807	FLAVANONE 3-HYDROXYLASE-LIKE PROTEIN	Hydroxylase
21945	2808	FLAVIN-CONTAINING MONOOXYGENASE, RHESUS	Oxygenases
		MACAQUE, SWISSPROT:FMO2_MACMU[PUTATIVE]	
21966	2809	RECEPTOR-LIKE PROTEIN KINASE - LIKE RECEPTOR-	Kinase, Protein
		LIKE PROTEIN KINASE 5, ARABIDOPSIS THALIANA,	
		PIR:S27756	
21974	2810	PROTEIN PHOSPHATASE - LIKE PROTEIN	Phosphatase
	,	PHOSPHATASE-2C, MESEMBRYANTHEMUM	
		CRYSTALLINUM, AF075579	

21987	2811	LYSINE DECARBOXYLASE, ARABIDOPSIS	Decarboxylase
		THALIANA[PUTATIVE]	
22004	2812	UTP-GLUCOSE GLUCOSYLTRANSFERASE - LIKE	Transferases
		PROTEIN UTP-GLUCOSE GLUCOSYLTRANSFERASE	,
		CASSAVA, PIR:S41951	
22010	2813	PYRUVATE WATER DIKINASE, ARCHAEOGLOBUS	Kinase
		FULGIDUS, PIR:F69338[PUTATIVE]	
22013	2814	ISOPENICILLIN N EPIMERASE, STREPTOMYCES	Epimerase
		CLAVULIGERUS, EMBL:M32324[PUTATIVE]	
22020	2815	PECTINACETYLESTERASE PRECURSOR - LIKE	Esterase
		PROTEIN PECTINACETYLESTERASE PRECURSOR,	
		VIGNA RADIATA, PIR:S68805	
22034	2816	PECTINESTERASE 2 PRECURSOR, ARABIDOPSIS	Esterase
		THALIANA, PIR:PC4168[PUTATIVE]	
22035	2817	FERROPORTINI, MUS MUSCULUS,	Transporter
		EMBL:AF226613[PUTATIVE]	
22041		RRNA METHYLASES[PUTATIVE]	Methylase
22047		DNA POLYMERASE SUBUNIT [PUTATIVE]	Polymerase
22063	2820	ION CHANNEL - LIKE PROTEIN LIGAND GATED	Channel
		CHANNEL-LIKE PROTEIN, BRASSICA NAPUS,	
22074		EMBL:AF109392	
22074		G protein SEVEN TRANSMEMBRANE DOMAIN ORPHAN	Receptor
		RECEPTOR, MUS MUSCULUS,	
22094		EMBL:AF051098[PUTATIVE] AMINO ACID AMINOTRANSFERASE BRANCHED-CHAIN	T
22094	- 1	AMINO ACID AMINOTRANSFERASE - PSEUDOMONAS	Transferases
	i	AERUGINOSA, SWISSPROT:ILVE PSEAE[PUTATIVE]	
22096		SIGNAL PEPTIDASE	Protease
22104			Kinase, Protein
	ĺ	KINASE - ARABIDOPSIS THALIANA, EMBL:U58918	Killase, Frotein
22106			Glycosylase
		POLYGALACTURONASE - LYCOPERSICON	, coo, 1000
	1	ESCULENTUM, EMBL:AF118567	
22132		PROTEIN KINASE - LIKE PROTEIN MEKK5 (ASKI,	Kinase, Protein
	I	MAPKKK5) GENE FOR MAP/ERK KINASE KINASE 5,	
		HOMO SAPIENS, TREMBL:HS325F22	
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WO 02/10210 PCT/EP01/09892

22140	2827	PECTIN METHYL-ESTERASE - LIKE PROTEIN PECTIN	Esterase
		  METHYL-ESTERASE PER, MEDICAGO TRUNCATULA,	}
		EMBL:MTR249611	ļ
22146	2828	PROTEIN PHOSPHATASE - LIKE PROTEIN	Phosphatase
		PHOSPHATASE-2C, MESEMBRYANTHEMUM	
		CRYSTALLINUM, EMBL:AF075581	
22189	2829	1-AMINOCYCLOPROPANE-1-CARBOXYLATE	Synthase
1		SYNTHASE 1-AMINOCYCLOPROPANE-1-	
		CARBOXYLATE SYNTHASE - ARABIDOPSIS THALIANA,	
		EMBL:U26542, FRAGMENT OF	
22221	2830	RECEPTOR-LIKE PROTEIN KINASE PRECURSOR - LIKE	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE, MADAGASCAR	
		PERIWINKLE, PIR:T10060	
22268	2831	GALACTINOL SYNTHASE - LIKE	Synthase
22340	2832	ADENYLATE KINASE -LIKE PROTEIN PREDICTED	Kinase
		PROTEINS, ARABIDOPSIS THALIANA	
22388	2833	RECEPTOR-LIKE PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE, ARABIDOPSIS	
		THALIANA, EMBL:ATRLPKB	
22390	2834	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE	Kinase, Protein
		PROTEIN KINASE PRK1, LYCOPERSICON	
		ESCULENTUM, PIR:T07865[PUTATIVE]	
22424	2835	BETA-1,3-GLUCANASE[PUTATIVE]	Glycosylase
22425	2836	HISTIDINE KINASE-LIKE PROTEIN	Kinase, Protein
22446	2837	SERINE/THREONINE PROTEIN KINASE-LIKE	Kinase, Protein
22475	2838	PROTEIN PHOSPHATASE-2C PP2C-LIKE	Phosphatase
22476	2839	ASPARTYL PROTEASE-LIKE	Protease
22477	2840	GSH-DEPENDENT DEHYDROASCORBATE REDUCTASE	Reductase
		1-LIKE	
22502	2841	GLUCAN SYNTHASE GLUCAN SYNTHASES -	Synthase
		DIFFERENT SPECIES[PUTATIVE]	
22521	2842	TRANSPORTER PROTEIN NA+/H+-EXCHANGING	Transporter
		PROTEIN NAPA - ENTEROCOCCUS HIRAE,	
		PIR:A42111[PUTATIVE]	j
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22532	2843	CAFFEIC ACID O-METHYLTRANSFERASE-LIKE Transferases
22332	2043	
		PROTEIN LIGNIN-BISPECIFIC O- METHYLTRANSFERASE - POPULUS TREMULOIDES,
		EMBL:X62096
22533	2844	SUCROSE SYNTHASE-LIKE PROTEIN SUCROSE Synthase
		SYNTHASE - CITRUS UNSHIU, EMBL:AB022091
22560	2845	RECEPTOR PROTEIN KINASE-LIKE PROTEIN  Kinase, Protein
22594	2846	PROTEIN KINASE - LIKE PROTEIN PTO KINASE Kinase, Protein
		INTERACTOR 1 (PTI1), LYCOPERSICON ESCULENTUM,
		EMBL:SL28007
22610	2847	GLUCURONOSYL TRANSFERASE-LIKE PROTEIN Transferases
		GLUCURONOSYL TRANSFERASE-LIKE PROTEIN,
		TOMATO, PIR:S39507
22611	2848	QUINONE OXIDOREDUCTASE (EC 1.6.5.5) P1, Reductase
		ARABIDOPSIS THALIANA, PIR:S57611[PUTATIVE]
22616	2849	GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
		GLUCOSE GLUCOSYLTRANSFERASE - SORGHUM
		BICOLOR, EMBL:AF199453
22617	2850	SAMT-LIKE PROTEIN S-ADENOSYL-L- Transferases
		METHIONINE:SALICYLIC ACID CARBOXYL
		METHYLTRANSFERASE (SAMT)- CLARKIA BREWERI,
		EMBL:AF133053
22619	2851	GLUCOSYLTRANSFERASE-LIKE PROTEIN UDP-Transferases
		GLUCOSE GLUCOSYLTRANSFERASE - SORGHUM
		BICOLOR, EMBL:AF199453
22628	2852	ANTHRANILATE N-BENZOYLTRANSFERASE -LIKE Transferases
		PROTEIN ANTHRANILATE N-BENZOYLTRANSFERASE
		(EC 2.3.1.144), CLOVE PINK, PIR:T10717
22636	2853	WALL-ASSOCIATED KINASE 4 (WAK4), ARABIDOPSIS Kinase, Protein
		THALIANA, EMBL:ATH9695
22639	2854	RECEPTOR SERINE/THREONINE PROTEIN KINASE - Kinase, Protein
		LIKE RECEPTOR SERINE/THREONINE KINASE PR5K,
		ARABIDOPSIS THALIANA, EMBL:AT48698
22640	2855	RECEPTOR SERINE/THREONINE PROTEIN KINASE - Kinase, Protein
		LIKE RECEPTOR SERINE/THREONINE KINASE PR5K,
		ARABIDOPSIS THALIANA, EMBL:AT48698
		-

22641	2856	RECEPTOR SERINE/THREONINE PROTEIN KINASE	- Kinase, Protein
		LIKE RECEPTOR SERINE/THREONINE KINASE PR5K	,
		ARABIDOPSIS THALIANA, EMBL:AT48698	
22656	2857	RIBULOSE BISPHOSPHATE CARBOXYLASE SMALL	Carboxylase
		CHAIN 3B PRECURSOR (RUBISCO SMALL SUBUNIT 3B	
		(SP P10798)	
22657	2858	RIBULOSE BISPHOSPHATE CARBOXYLASE SMALL	Carboxylase
		CHAIN 2B PRECURSOR (RUBISCO SMALL SUBUNIT 2B)	
		(SP P10797)	
22658	2859	RIBULOSE BISPHOSPHATE CARBOXYLASE SMALL	Carboxylase
		CHAIN 1B PRECURSOR (RUBISCO SMALL SUBUNIT 1B)	)
		(SP P10796)	<u></u>
22668		TRYPTOPHAN SYNTHASE BETA CHAIN	Synthase
22671	2861	PROTEIN KINASE[PUTATIVE]	Kinase, Protein
22697	2862	TRANSPORTER -LIKE PROTEIN N SYSTEM AMINO	Transporter
		ACIDS TRANSPORTER NAT-1, MUS MUSCULUS,	
		EMBL:AF159856	
22717	2863	RECEPTOR PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
İ		RECEPTOR-PROTEIN KINASE-LIKE PROTEIN,	
		ARABIDOPSIS THALIANA, PIR:T45786	
22718	2864	RECEPTOR PROTEIN KINASE - LIKE PROTEIN	Kinase, Protein
		RECEPTOR SERINE/THREONINE KINASE PR5K,	
		ARABIDOPSIS THALIANA, EMBL:AT48698	
22720	2865	ACYLTRANSFERASE - LIKE PROTEIN ANTHOCYANIN 5-	Transferases
		AROMATIC ACYLTRANSFERASE, GENTIANA	
		TRIFLORA, EMBL:AB010708	
22723	2866	ACYLTRANSFERASE -LIKE PROTEIN ANTHOCYANIN 5-	
		AROMATIC ACYLTRANSFERASE, GENTIANA	
2272 (	2065	TRIFLORA, EMBL: AB010708	
22724		ACYLTRANSFERASE -LIKE PROTEIN ANTHOCYANIN	Transferases
	- 1	ACYLTRANSFERASE, PERILLA FRUTESCENS,	
22754		EMBL:AB029340	
22754		RECEPTOR PROTEIN KINASE -LIKE PROTEIN PROTEIN	Kinase, Protein
22772		KINASE XA21, RICE, PIR:A57676	
22773	2869	PEROXIDASE ATP24A	Oxidase

22798	2870	DEGP PROTEASE-LIKE PROTEIN DEGP PROTEASE	Protease
		PRECURSOR - ARABIDOPSIS THALIANA,	ļ
		EMBL:AF028842	·
22806	2871	POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
		POLYGALACTURONASE PRECURSOR - LYCOPERSICON	
		ESCULENTUM, PIR:S57806	
22830	2872	PEROXIDASE ATP26A	Oxidase
22835	2873	PROTEASE SERINE PROTEINASE DO, BACILLUS	Protease
		SUBTILIS, PIR:A69643[PUTATIVE]	
22853	2874	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE	Kinase, Protein
		SERINE/THREONINE KINASE RKF2, ARABIDOPSIS	
		THALIANA, EMBL:AF024649	
22854	2875	RAFFINOSE SYNTHASE -LIKE PROTEIN RAFFINOSE	Synthase
		SYNTHASE RFS, CUCUMIS SATIVUS, EMBL:AF073744	
22893	2876	AMINO ACID PERMEASE	Transporter
22900	2877	UROPHORPHYRIN III METHYLASE (GB AAB92676.1)	Methylase
22914	2878	LIPASE/HYDROLASE GDSL-like -motif	Lipase
22919	2879	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE-	Transferases
	٠	LIKE PROTEIN	
22931	2880	PURINE PERMEASE-LIKE PROTEIN	Transporter
22933	2881	RECEPTOR KINASE-LIKE PROTEIN	Kinase, Protein
22937	2882	GLUTATHIONE TRANSFERASE-LIKE	Transferases
22939	2883	GLUTATHIONE TRANSFERASE	Transferases
22941	2884	PROTEIN KINASE-LIKE	Kinase, Protein
22962	2885	FOLYLPOLYGLUTAMATE SYNTHASE-LIKE PROTEIN	Synthase
22975	2886	NA+/H+ ANTIPORTER-LIKE PROTEIN	Transporter
22990	2887	CMP-SIALIC ACID TRANSPORTER-LIKE PROTEIN	Transporter
22994	2888	AMINO ACID PERMEASE-LIKE PROTEIN; PROLINE	Transporter
		TRANSPORTER-LIKE PROTEIN	- {
23001	2889	POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
23003	2890	LIPASE/HYDROLASE GDSL-like -motif	Lipase
23024	2891	BETA-1,3-GLUCANASE-LIKE PROTEIN	Glycosylase
23026	2892	RECEPTOR LECTIN KINASE-LIKE PROTEIN	Kinase, Protein
23031	2893	LIPASE/HYDROLASE GDSL-like -motif	Lipase

23032		PEROXIDASE (EMB CAA66960.1)	Oxidase
23035	2895	TETRACYCLINE TRANSPORTER PROTEIN[PUTATIVE]	Transporter
23058	2896	SERINE/THREONINE PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
23079	2897	ALLENE OXIDE SYNTHASE (EMB CAA73184.1)	Synthase
23086	2898	BETA-1,3-GLUCANASE-LIKE PROTEIN	Glycosylase
23094	2899	DIHYDROFLAVONOL 4-REDUCTASE	Reductase
23097	2900	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE-	Transferases
		LIKE PROTEIN	-
23116	2901	RECEPTOR KINASE-LIKE PROTEIN	Kinase, Protein
23148	2902	INORGANIC PHOSPHATE TRANSPORTER	Transporter
1.		(DBJ BAA34390.1)	
23149	2903	PHOSPHATE TRANSPORTER (GB AAB17265.1)	Transporter
23150	2904	INORGANIC PHOSPHATE TRANSPORTER	Transporter
		(DBJ BAA24281.1)	-
23151	2905	INORGANIC PHOSPHATE TRANSPORTER	Transporter
<u> </u>		(DBJ BAA24282.1)	
23157	2906	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
23158	2907	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE	Oxidase
23173	2908	N-CARBAMYL-L-AMINO ACID AMIDOHYDROLASE-	Hydrolase
		LIKE PROTEIN	
23174		SUCROSE TRANSPORTER PROTEIN	Transporter
23189		BETA-KETOACYL-COA SYNTHASE	Synthase
23202		DIMETHYLANILINE MONOOXYGENASE-LIKE	Oxygenases
23216	2912	CELLULOSE SYNTHASE CATALYTIC SUBUNIT-LIKE	Synthase
		PROTEIN	
23265		RIBOSE 5-PHOSPHATE ISOMERASE[PUTATIVE]	Isomerase
23276		TERPENE CYCLASE/SYNTHASE	Cyclase
23281			Glycosylase
23283	2916	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
23296	2917	POLYGALACTURONASE-LIKE PROTEIN	Glycosylase
23297	2918	POLYGALACTURONASE	Glycosylase
23331	2919	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE-	Oxygenases
		FORMING)-LIKE PROTEIN	j
23341	2920	PECTINACETYLESTERASE	Esterase
23343	2921	BETA-AMYLASE-LIKE	Glycosylase

23377 2923 SUBTILISIN-LIKE PROTEASE Protease 23378 2924 SUBTILISIN-LIKE PROTEASE Protease 23379 2925 LIPASE/HYDROLASE GDSL-like -motif Lipase 23380 2926 PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-TYPE Isomera [PUTATIVE] 23384 2927 DNA POLYMERASE III, GAMMA SUBUNIT[PUTATIVE] Polymera 23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase, 23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN 23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase, 23396 2931 RECEPTOR PROTEIN KINASE-LIKE PROTEIN Kinase,	rase Protein
23379 2925 LIPASE/HYDROLASE GDSL-like -motif Lipase 23380 2926 PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-TYPE Isomera [PUTATIVE] 23384 2927 DNA POLYMERASE III, GAMMA SUBUNIT[PUTATIVE] Polymera 23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase, 23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN 23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	rase Protein
23380 2926 PEPTIDYL-PROLYL CIS-TRANS ISOMERASE FKBP-TYPE Isomera [PUTATIVE]  23384 2927 DNA POLYMERASE III, GAMMA SUBUNIT[PUTATIVE] Polymer 23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase, PROTEIN  23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN  23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase, Kinase,	rase Protein atase
[PUTATIVE]  23384 2927 DNA POLYMERASE III, GAMMA SUBUNIT[PUTATIVE] Polymer  23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase,  23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN  23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	Protein ntase
23384 2927 DNA POLYMERASE III, GAMMA SUBUNIT[PUTATIVE] Polymer 23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase, 23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN 23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	Protein
23390 2928 RECEPTOR-LIKE PROTEIN KINASE Kinase,  23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN  23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	Protein
23391 2929 TYROSINE-SPECIFIC PROTEIN PHOSPHATASE-LIKE Phospha PROTEIN  23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	ntase
PROTEIN  23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	
23392 2930 RECEPTOR KINASE-LIKE PROTEIN Kinase,	Protein
	Protein
23396 2931 RECEPTOR PROTEIN KINASE-LIKE PROTEIN Kinase	i ioiciii
I I I I I I I I I I I I I I I I I I I	Protein
23403 2932 LIPASE/HYDROLASE GDSL-like -motif Lipase	
23405 2933 MAGNESIUM CHELATASE SUBUNIT OF Chelatas	ie .
PROTOCHLOROPHYLLIDE REDUCTASE	
23407 2934 LIPASE/HYDROLASE GDSL-like -motif Lipase	
23408 2935 LIPASE/HYDROLASE GDSL-like -motif Lipase	
23420 2936 SERINE/THREONINE PROTEIN KINASE-LIKE PROTEIN Kinase,	Protein
23449 2937 POTASSIUM CHANNEL OUTWARD RECTIFYING KCO Channel	
23467 2938 KINASE[PUTATIVE] Kinase,	Protein
23469 2939 PROTEIN KINASE-LIKE PROTEIN Kinase,	Protein
23511 2940 PEROXIDASE Oxidase	
23516 2941 S-RIBONUCLEASE BINDING PROTEIN[PUTATIVE] Nuclease	<del></del>
23518 2942 SERINE/THREONINE KINASE-LIKE Kinase,	Protein
23556 2943 MEMBRANE CHANNEL PROTEIN-LIKE; AQUAPORIN Channel	
(TONOPLAST INTRINSIC PROTEIN)-LIKE	
23561 2944 PECTIN METHYLESTERASE-LIKE Esterase	~
23592 2945 PYROPHOSPHATE-FRUCTOSE-6-PHOSPHATE 1- Transfer	ases
PHOSPHOTRANSFERASE-LIKE PROTEIN	
23595 2946 ADENYLATE KINASE[PUTATIVE] Kinase	
23617 2947 PHOSPHORIBOSYLANTHRANILATE TRANSFERASE-Transfer	ases
LIKE PROTEIN	
23618 2948 XYLOGLUCAN ENDO-1,4-BETA-D-GLUCANASE Glycosyl	350

		<u> </u>	
23621		LACCASE (DIPHENOL OXIDASE)	Oxidase
23622	<u> </u>	TERPENE SYNTHASE	Synthase
23625		POLYGALACTURONASE	Glycosylase
23633	2952	INDOLE-3-GLYCEROL PHOSPHATE SYNTHASE	Synthase
23649	2953	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
23651	2954	LIGAND-GATED ION CHANNEL PROTEIN-LIKE	Channel
		GLUTAMATE RECEPTOR-LIKE	
23652	2955	LIGAND-GATED ION CHANNEL PROTEIN-LIKE	Channel
		GLUTAMATE RECEPTOR-LIKE	
23656	2956	PECTINESTERASE-LIKE; ALSO HIGHLY SIMILAR TO L-	Oxidase
		ASCORBATE OXIDASE AND POLLEN-SPECIFIC	
		PROTEIN[PUTATIVE]	
23685	2957	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
23695	2958	PANTOATE-BETA-ALANINE LIGASE	Ligase
23701	2959	PECTATE LYASE	Lyase
23704	2960	ANTHRANILATE N-BENZOYLTRANSFERASE	Transferases
23718	2961	FATTY ACID ELONGASE; BETA-KETOACYL-COA	Synthase
		SYNTHASE-LIKE PROTEIN	
23729	2962	PECTIN METHYLESTERASE	Esterase
23730	2963	SUCROSE SYNTHASE	Synthase
23738	2964	PHYTOCHELATIN SYNTHETASE[PUTATIVE], PROTEIN	Synthase
		CONTAINS SIMILARITY TO	
23747	2965	XYLOSIDASE	Glycosylase
23774	2966	AMINO ACID PERMEASE 6 (EMB CAA65051.1)	Transporter
23777	2967	RECEPTOR PROTEIN KINASE	Kinase, Protein
23780	2968	ANTHOCYANIDIN-3-GLUCOSIDE	Transferases
		RHAMNOSYLTRANSFERASE-LIKE	]
23784	2969	FRO2-LIKE PROTEIN; NADPH OXIDASE-LIKE	Oxidase
23785	2970	FROI-LIKE PROTEIN; NADPH OXIDASE-LIKE	Oxidase
23786	2971	ELICITOR-INDUCIBLE RECEPTOR-LIKE PROTEIN	Receptor
]		EIR[PUTATIVE]	-
23787	2972	RECEPTOR PROTEIN KINASE-LIKE	Kinase, Protein
22705	2070	DECEDED DO COMPOSITOR AND A CONTROL OF A CON	
23788	29/3	RECEPTOR PROTEIN KINASE-LIKE	Kinase, Protein

23789	2974	RECEPTOR PROTEIN KINASE-LIKE	Kinase, Protein
23792	2975	METHIONINE S-METHYLTRANSFERASE	Transferases
		(GB AAD49574.1)	
23844	2976	ABC TRANSPORTER[PUTATIVE]	Transporter
23870	2977	11-BETA-HYDROXYSTEROID DEHYDROGENASE-LIKE	Dehydrogenases
23871	2978	11-BETA-HYDROXYSTEROID DEHYDROGENASE-LIKE	Dehydrogenases
23876	2979	PROTEIN TRANSPORT PROTEIN SEC12P-LIKE	Transporter
23880	2980	11-BETA-HYDROXYSTEROID DEHYDROGENASE-LIKE	Dehydrogenases
23881	2981	11-BETA-HYDROXYSTEROID DEHYDROGENASE-LIKE	Dehydrogenases
23886	2982	UDP-GLUCOSE:PROTEIN TRANSGLUCOSYLASE;	Glycosylase
		REVERSIBLY GLYCOSYLATED POLYPEPTIDE	
23888	2983	OXIDOREDUCTASE[PUTATIVE]	Reductase
23937	2984	ACID PHOSPHATASE	Phosphatase
23942	2985	GIBBERELLIN 20-OXIDASE-LIKE PROTEIN	Oxidase
23946	2986	RECEPTOR PROTEIN KINASE-LIKE	Kinase, Protein
23953	2987	WAX SYNTHASE[PUTATIVE], PROTEIN CONTAINS SIMILARITY TO	Synthase
23956		AUTOCRINE MOTILITY FACTOR RECEPTOR[PUTATIVE]	Receptor
23957	2989	TREHALOSE-6-PHOSPHATE PHOSPHATASE	Phosphatase
23959	2990	PECTINESTERASE-LIKE; STRONG SIMILARITY TO	Esterase
		POLLEN-SPECIFIC PROTEIN[PUTATIVE]	
23960	2991	PECTINESTERASE	Esterase
23961	2992	PECTINESTERASE	Esterase
23967	2993	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
23982	2994	POTASSIUM/PROTON ANTIPORTER-LIKE PROTEIN	Transporter
23985	2995	PEPTIDASE[PUTATIVE]	Protease
23986	2996	SERINE PROTEASE-LIKE PROTEIN	Protease
23987	2997	PROTEIN PHOSPHATASE-2C; PP2C-LIKE PROTEIN	Phosphatase
23988	2998	PROTEIN KINASE[PUTATIVE]	Kinase, Protein
23992	2999	GIBBERELLIN 20-OXIDASE (EMB CAA58294.1)	Oxidase

PCT/EP01/09892

23994	3000	FRUCTOKINASE 1	Kinase
24000	3001	PEROXIDASE	Oxidase
24004	3002	MANDELONITRILE LYASE-LIKE PROTEIN	Lyase
24009	3003	MYOSIN HEAVY CHAIN KINASE[PUTATIVE],	Kinase, Protein
		CONTAINS SIMILARITY TO	ł
24021	3004	DIHYDRODIPICOLINATE REDUCTASE-LIKE PROTEIN	Reductase
24039	3005	HYALURONAN MEDIATED MOTILITY RECEPTOR-LIKE	Receptor
		PROTEIN	
24068		BETA-CAROTENE HYDROXYLASE	Hydroxylase
24091	3007	ORNITHINE CYCLODEAMINASE[PUTATIVE PROTEIN	Deaminase
		CONTAINS SIMILARITY TO]	
24119	3008	PROTOCHLOROPHYLLIDE REDUCTASE;	ſ
		OXIDOREDUCTASE REQUIRED FOR SHOOT APEX	
		DEVELOPMENT	
24120	3009	PROTOCHLOROPHYLLIDE REDUCTASE;	
		OXIDOREDUCTASE REQUIRED FOR SHOOT APEX	
		DEVELOPMENT	
24123	3010	CYCLIC NUCLEOTIDE-REGULATED ION CHANNEL	Channel
		(EMB CAA76178.1)	
24142	3011	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
24147	3012	PECTINESTERASE	Esterase
24168	3013	ALDO/KETO REDUCTASE-LIKE PROTEIN	Reductase
24191	3014	CAFFEIC ACID 3-O-METHYLTRANSFERASE-LIKE	Transferases
		PROTEIN	
24199	3015	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
24202	3016	METHYLTRANSFERASE-LIKE PROTEIN , RIBOSOMAL	Transferases
		PROTEIN L11	,
24209	3017	FLAVONOL 3-O-GLUCOSYLTRANSFERASE-LIKE	Transferases
		PROTEIN	ļ
24210	3018	ETHYLENE-FORMING-ENZYME-LIKE DIOXYGENASE	Oxygenases
24211	3019	FLAVONOL 3-O-GLUCOSYLTRANSFERASE-LIKE	Transferases
		PROTEIN	ł
24216	3020	FLAVONOL 3-O-GLUCOSYLTRANSFERASE-LIKE	Transferases
24224	3021	IAA-AMINO ACID HYDROLASE HOMOLOG ILL3	Hydrolase
	j	(GB AAC31939.1)	,
	· · · · · · · · · · · · · · · · · · ·		

24226	3022	O-METHYLTRANSFERASE	Transferases
24229	3023	PROTOCHLOROPHYLLIDE OXIDOREDUCTASE A	Reductase
		NADPH (GB AAC49043.1)	
24235	3024	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
2425	302-	ION CHANNEL	
24248	2025	RECEPTOR-PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
24246	3023	RECEI TOR-I ROTEIN KINASE-LIKE I ROTEIN	Kinase, Flotein
24260	3026	1,4-BENZOQUINONE REDUCTASE-LIKE; TRP	Reductase
	j	REPRESSOR BINDING PROTEIN-LIKE	
24269	3027	SERINE/THREONINE-SPECIFIC PROTEIN KINASE-LIKE	Kinase, Protein
		PROTEIN	
24291	3028	TRYPTOPHAN SYNTHASE BETA CHAIN 1 PRECURSOR	Synthase
		(SP P14671)	
24306	3029	PYRUVATE DECARBOXYLASE (GB AAB16855.1)	Decarboxylase
24315	3030	LIPASE/HYDROLASE GDSL-like -motif	Lipase
24328		BETA-1,3-GLUCANASE-LIKE PROTEIN	Glycosylase
24335			Transferases
		CARBOXYL METHYLTRANSFERASE-LIKE PROTEIN	
24342	3033	WAX SYNTHASE-LIKE PROTEIN	Synthase
24343	3034	WAX SYNTHASE-LIKE PROTEIN	Synthase
24344	3035	WAX SYNTHASE[PUTATIVE], PROTEIN CONTAINS	Synthase
		SIMILARITY TO	
24345	3036	WAX SYNTHASE-LIKE PROTEIN	Synthase
24346	3037	WAX SYNTHASE-LIKE PROTEIN	Synthase
24347	3038	WAX SYNTHASE-LIKE PROTEIN	Synthase
24348	3039	WAX SYNTHASE-LIKE PROTEIN	Synthase
24369	3040	PECTIN METHYLESTERASE [PUTATIVE]	Esterase
24373	3041	POTASSIUM CHANNEL OUTWARD RECTIFYING KCO	Channel
24380	3042	BETA-AMYLASE	Glycosylase
24382	3043	PECTATE LYASE	Lyase
24393	3044	SERINE/THREONINE-SPECIFIC KINASE LIKE PROTEIN	Kinase, Protein
24414	3045	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
24418	3046	NICOTIANAMINE SYNTHASE	Synthase

24436	3047	S-ADENOSYLMETHIONINE:2-	Transferases
		DEMETHYLMENAQUINONE METHYLTRANSFERASE	1
		LIKE	
24440	3048	S-ADENOSYL-L-METHIONINE:SALICYLIC ACID	Transferases
		CARBOXYL METHYLTRANSFERASE-LIKE PROTEIN	
24443	3049	CARBONIC ANHYDRASE[PUTATIVE, PROTEIN	Anhydrase
		CONTAINS SIMILARITY TO]	
24469	3050	BETA-1,3-GLUCANASE-LIKE PROTEIN	Glycosylase
24473	3051	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE-	Kinase
		DEPENDENT PHOSPHOFRUCTO-1-KINASE-LIKE	,
		PROTEIN	•
24475	3052	1	Hydrolase
		PRECURSOR (SP P54969)	
24476		IAA-AMINO ACID HYDROLASE (GB AAC04866.1)	Hydrolase
24483		ZINC PROTEASE PQQL-LIKE PROTEIN	Protease
24484	3055	HISTONE ACETYLTRANSFERASE HAT B	Transferases
24486	3056	SERINE O-ACETYLTRANSFERASE (EC 2.3.1.30) SAT-52	Transferases
		(PIR  S71207)	
24489	3057	PROTEIN KINASE[PUTATIVE]	Kinase, Protein
24506	3058	CYTOKININ OXIDASE	Oxidase
24512	3059	LYCOPENE EPSILON CYCLASE	Cyclase
24518	3060	AUXIN TRANSPORT PROTEIN EIR1 (GB AAC39513.1)	Transporter
24528	3061	PHOSPHATIDYLSERINE DECARBOXYLASE	Decarboxylase
24555	3062	AAA-TYPE ATPASE[PUTATIVE]	ATPase
24560	3063	XYLOGLUCAN ENDOTRANSGLYCOSYLASE	Glycosylase
24561	3064	XYLOGLUCAN ENDOTRANSGLYCOSYLASE	Glycosylase
24562	3065	ENDOXYLOGLUCAN TRANSFERASE (GB AAD45127.1)	Transferases
24566	3066	ADENOSYLMETHIONINE-8-AMINO-7-OXONONANOATE	Transferases
		AMINOTRANSFERASE-LIKE PROTEIN	
24591	3067	N-HYDROXYCINNAMOYL/BENZOYLTRANSFERASE	Transferases
24592	3068	BRANCHED-CHAIN AMINO ACID	Transferases
		AMINOTRANSFERASE-LIKE PROTEIN	
24596	3069	ANTHRANILATE SYNTHASE BETA CHAIN	Synthase

		T	
24601	3070	CYCLIC NUCLEOTIDE AND CALMODULIN-REGULATED	Channel
		ION CHANNEL (EMB CAB40130.1)	
24604	3071	DNA-3-METHYLADENINE GLYCOSYLASE[PUTATIVE]	Glycosylase
24616	3072	BETA-1,3 GLUCANASE - LIKE PROTEIN BETA-1,3	Glycosylase
i	<u> </u>	GLUCANASE, POPULUS ALBA X POPULUS TREMULA,	1
		EMBL:AF230109	
24622	3073	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
24627	3074	PHOSPHOESTERASE [CONTAINS SIMILARITY TO]	Esterase
24637	3075	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
24638	3076	ALPHA-HYDROXYNITRILE LYASE, POLYNEURIDINE	Lypac
24038	3070	ALDEHYDE ESTERASE-LIKE; ALSO SIMILAR TO	
		ALPHA-HYDROXYNITRILE LYASE	
24646	2077	PEROXIDASE	Oxidase
	İ	PEROXIDASE	
24647	·		Oxidase
24655	3079	BETA 1-3 GLUCANASE - LIKE PROTEIN BETA 1-3	Glycosylase
24656	2000	GLUCANASE, VITIS VINIFERA, EMBL:VVI277900	
24656	3080	CINNAMOYL-COA REDUCTASE - LIKE PROTEIN	Reductase
!		CINNAMOYL-COA REDUCTASE, CIDER TREE,	
21550		PIR:T10733	
24659	1	PROTEIN KINASE - LIKE PROTEIN KINASE ATNI,	Kinase, Protein
		ARABIDOPSIS THALIANA, PIR:S61766	
24661			Kinase, Protein
		NPK15, NICOTIANA TABACUM, PIR:S52578[PUTATIVE]	
24680		CARBOHYDRATE KINASE, PFKB, ARCHAEOGLOBUS	Kinase
		FULGIDUS, PIR:A69300[PUTATIVE]	
24701		RECEPTOR-LIKE PROTEIN KINASE PRECURSOR - LIKE	Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE PRECURSOR,	
		MADAGASCAR PERIWINKLE, PIR:T10060	
24708	3085		Kinase, Protein
		SERINE/THREONINE/TYROSINE-SPECIFIC PROTEIN	
		KINASE APK1, ARABIDOPSIS THALIANA, PIR:S28615	
24710	3086	COPPER TRANSPORT PROTEIN	Transporter
24711	3087	COPPER TRANSPORT PROTEIN - LIKE	Transporter
24714	3088	HEXOSYLTRANSFERASE - LIKE PROTEIN	Transferases

24733	3089	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE Kinase, Protein
ĺ		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		PIR:T47481
24734	3090	RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE Kinase, Protein
		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		PIR:T47481
24759	3091	ZINC TRANSPORTER ZIP2 - LIKE Z25114[PUTATIVE] Transporter
24760	3092	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE - Oxidase
1		LIKE PROTEIN 1-AMINOCYCLOPROPANE-1-
		CARBOXYLATE OXIDASE KIDNEY BEAN, PIR:T10818
24761	3093	1-AMINOCYCLOPROPANE-1-CARBOXYLATE OXIDASE - Oxidase
		LIKE PROTEIN 1-AMINOCYCLOPROPANE-1-
		CARBOXYLATE OXIDASE KIDNEY BEAN, PIR:T10818
24765	3094	GLUCURONOSYL TRANSFERASE - LIKE PROTEIN Transferases
		GLUCURONOSYL TRANSFERASE HOMOLOG,
		RIPENING-RELATED, LYCOPERSICON ESCULENTUM,
		PIR:S39507
24766	3095	GLUCURONOSYL TRANSFERASE - LIKE PROTEIN Transferases
		GLUCURONOSYL TRANSFERASE HOMOLOG,
		RIPENING-RELATED, LYCOPERSICON ESCULENTUM,
		PIR:S39507
24771	3096	SERINE/THREONINE-SPECIFIC PROTEIN KINASE - LIKE Kinase, Protein
		PUTATIVE PROTEIN SERINE /THREONINE KINASE,
		SORGHUM BICOLOR, EMBL:SBRLK1
24772	3097	SERINE/THREONINE-SPECIFIC PROTEIN KINASE - LIKE Kinase, Protein
		PUTATIVE PROTEIN SERINE /THREONINE KINASE,
		SORGHUM BICOLOR, EMBL:SBRLK1
24773	3098	SERINE/THREONINE-SPECIFIC PROTEIN KINASE - LIKE Kinase, Protein
		PUTATIVE PROTEIN SERINE /THREONINE KINASE,
0.455.4	2000	SORGHUM BICOLOR, EMBL:SBRLK1
24774		SERINE/THREONINE-SPECIFIC PROTEIN KINASE - LIKE Kinase, Protein
		PUTATIVE PROTEIN SERINE /THREONINE KINASE,
0.177	2100	SORGHUM BICOLOR, EMBL:SBRLK1
24777	3100	RECEPTOR-LIKE PROTEIN KINASE PRECURSOR - LIKE Kinase, Protein
		RECEPTOR-LIKE PROTEIN KINASE PRECURSOR,
[		MADAGASCAR PERIWINKLE, PIR:T10060

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24781	3101	SERINE /THREONINE KINASE - LIKE PUTATIVE SERINE Kinase, Protein
		/THREONINE KINASE, SORGHUM BICOLOR,
		EMBL:SBRLK1
24782	3102	GTP CYCLOHYDROLASE II / 3,4-DIHYDROXY-2-Synthase
		BUTANONE-4-PHOSHATE SYNTHASE - LIKE PROTEIN
		GTP CYCLOHYDROLASE II / 3,4-DIHYDROXY-2-
		BUTANONE-4-PHOSHATE SYNTHASE, ARABIDOPSIS
		THALIANA, EMBL:ATAJ0053
24784	3103	TYROSINE PHOSPHATASE-LIKE PROTEIN, PTPLB, MUS Phosphatase
		MUSCULUS, EMBL:AF169286[PUTATIVE]
24788	3104	SUBTILISIN-LIKE PROTEASE - LIKE PROTEIN Protease
		SUBTILISIN-LIKE PROTEASE AIR3, ARABIDOPSIS
1 1		THALIANA, EMBL:AF098632
24815	3105	PROTEIN KINASE - LIKE PROTEIN KINASE 1, POPULUS Kinase, Protein
		NIGRA, EMBL:AB041503
24816	3106	PROTEIN KINASE - LIKE PROTEIN KINASE 1, POPULUS Kinase, Protein
		NIGRA, EMBL:AB041503
24830	3107	TRNA INTRON ENDONUCLEASE - LIKE PROTEIN TRNA Nuclease
		INTRON ENDONUCLEASE, ARABIDOPSIS THALIANA,
		EMBL:AB036339
24834	3108	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		PIR:T47484
24835	3109	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		PIR:T47484
24837	3110	RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
]		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		EMBL:AL138657
24838		RECEPTOR LIKE PROTEIN KINASE RECEPTOR LIKE Kinase, Protein
= .52.5		PROTEIN KINASE, ARABIDOPSIS THALIANA,
		EMBL:AL138657
24839		RECEPTOR-LIKE PROTEIN KINASE RECEPTOR-LIKE Kinase, Protein
24039		
		1
		PIR:T47473

24861	3113	IMIDAZOLEGLYCEROL-PHOSPHATE SYNTHASE Synthase
		SUBUNIT H - LIKE IMIDAZOLEGLYCEROL-PHOSPHATE
		SYNTHASE SUBUNIT H HOMOLOG, ARCHAEOGLOBUS
		FULGIDUS, PIR:E69313
24884	3114	HIGH-AFFINITY NITRATE TRANSPORTER ACHI - LIKE Transporter
		PROTEIN HIGH-AFFINITY NITRATE TRANSPORTER
		ACHI, ARABIDOPSIS THALIANA, EMBL:AF019748
24885	3115	HIGH AFFINITY NITRATE TRANSPORTER PROTEIN - Transporter
		LIKE PROBABLE HIGH AFFINITY NITRATE
		TRANSPORTER PROTEIN, GLYCINE MAX, PIR:T06237
24897	3116	S-RECEPTOR KINASE HOMOLOG 2 PRECURSOR S-Kinase, Protein
		RECEPTOR KINASE HOMOLOG 2 PRECURSOR,
		ARABIDOPSIS THALIANA, PIR:S27754
24899	3117	PHYTOCHELATIN SYNTHETASE - LIKE PROTEIN Synthase
		PUTATIVE PHYTOCHELATIN SYNTHETASE,
		ARABIDOPSIS THALIANA, EMBL:ATH6787
24902	3118	PHYTOCHELATIN SYNTHETASE - LIKE PUTATIVE Synthase
		PHYTOCHELATIN SYNTHETASE, ARABIDOPSIS
0.4000	2110	THALIANA, EMBL:ATH6787[PUTATIVE]
24923	3119	ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE - Transferases
		LIKE PROTEIN ANTHOCYANIN 5-AROMATIC ACYLTRANSFERASE, GENTIANA TRIFLORA,
		EMBL:AB010708
24942	3120	RECEPTOR-LIKE PROTEIN KINASE PRECURSOR - LIKE Kinase, Protein
24742	3120	RECEPTOR-LIKE PROTEIN KINASE PRECURSOR,
		MADAGASCAR PERIWINKLE, PIR:T10060
24955	3121	LEUCINE-RICH RECEPTOR-LIKE PROTEIN KINASE - Kinase, Protein
		LIKE PROTEIN LEUCINE-RICH RECEPTOR-LIKE
		PROTEIN KINASE LRPKMI, MALUS DOMESTICA,
		EMBL:AF053127
24964	3122	PROTEIN KINASE-LIKE TRANSMEMBRANE PROTEIN Kinase, Protein
		TMKLI PRECURSOR, ARABIDOPSIS THALIANA,
		EMBL:ATTMKL1[PUTATIVE]
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0.40.65	2100	The second secon	T
24965	3123	PHOSPHOFRUCTO-1-KINASE, PYROPHOSPHATE	İ
·		DEPENDENT PHOSPHOFRUCTO-1-KINASE - LIKE	1
		PROTEIN PYROPHOSPHATE-DEPENDENT	į
		PHOSPHOFRUCTO-1-KINASE, PRUNUS ARMENIACA	,
		EMBL:U93272	1
24975	3124	PECTIN METHYL-ESTERASE - LIKE PROTEIN PECTIN	Esterase
		METHYL ESTERASE PEST2, SOLANUM TUBEROSUM	
] ]		EMBL:AF152172	
25021	3125	IRON-REGULATED TRANSPORTER - LIKE PROTEIN	Transporter
		IRON-REGULATED TRANSPORTER 1, LYCOPERSICON	
1		ESCULENTUM, EMBL:AF136579	]
25028	3126	RECEPTOR-LIKE PROTEIN KINASE 5, ARABIDOPSIS	Kinase, Protein
		THALIANA, PIR:S27756[PUTATIVE]	
25053	3127	GLUTATHIONE S-TRANSFERASE-LIKE PROTEIN	Transferases
25058	3128	DEHYDROGENASE	Dehydrogenases
25065	3129	TRANSPORTIN-SR[PUTATIVE]	Transporter
25073	3130	PEPTIDE TRANSPORTER	Transporter
25076	3131	RECEPTOR - LIKE PROTEIN KINASE - LIKE PROTEIN	]
ĺ		SERINE/THREONINE KINASE RLK1, SORGHUM	,
		BICOLOR, EMBL:SBRLK1	
25078	3132	NITRATE TRANSPORTER NTLI - LIKE PROTEIN	Transporter
}		NITRATE TRANSPORTER NTL1, ARABIDOPSIS	
		THALIANA, EMBL:AF073361	
25084		1-DEOXY-D-XYLULOSE 5-PHOSPHATE	Isomerase
2500-1		REDUCTOISOMERASE (DXR)	Isomerase
25102		` '	Aldolase
25102			
			Lipase
25122			Lyase
25127			Glycosylase
25137	3138	KINASE[PUTATIVE]	Kinase, Protein
25145	3139	RECEPTOR-LIKE PROTEIN KINASE[PUTATIVE] Kinase, Protein	
25155	3140	ACETYLTRANSFERASE[PUTATIVE] Transferases	
25160	3141	ACYLTRANSFERASE-LIKE PROTEIN	Transferases

25161	3142	GLUTAMATE-1-SEMIALDEHYDE 2,1-AMINOMUTASE 1	
		PRECURSOR (GSA 1) (GLUTAMATE-1-SEMIALDEHYDE	
		AMINOTRANSFERASE 1) (GSA-AT 1) (SP P42799)	
25162	3143	FLAVONOL SYNTHASE	Synthase
25163	3144	FLAVONOL SYNTHASE	Synthase
25164	3145	1-AMINOCYCLOPROPANE-1-CARBOXYLIC ACID	Oxidase
		OXIDASE-LIKE PROTEIN	
25175	3146	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
25185	3147	BETA-GALACTOSIDASE (EMB CAB64746.1)	Glycosylase
25189	3148	AMINO ACID TRANSPORTER AAP4 (PIR  S51169)	Transporter
25193	3149	HISTIDINOL DEHYDROGENASE	Dehydrogenases
25197	3150	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
25198	3151	PROTEIN KINASE[PUTATIVE]	Kinase, Protein
25214	3152	PEROXIDASE ATP3A (EMB CAA67340.1)	Oxidase
25215	3153	PEROXIDASE ATP3A HOMOLOG	Oxidase
25216	3154	PEROXIDASE (EMB CAA67551.1)	Oxidase
25225	3155	ALTERNATIVE OXIDASE 2 (SP O22049)	Oxidase
25229	3156	2-NITROPROPANE DIOXYGENASE-LIKE PROTEIN	Oxygenases
25234	3157	GTP CYCLOHYDROLASE II; 3,4-DIHYDROXY-2-	Synthase
		BUTANONE-4-PHOSHATE SYNTHASE	•
	-	(EMB CAA03884.1)	
25261	3158	BETA-XYLOSIDASE	Glycosylase
25268	3159	PECTIN METHYLESTERASE-LIKE PROTEIN	Esterase
25278	3160	CELLULOSE SYNTHASE CATALYTIC SUBUNIT	Synthase
25283	3161	BETA-1,3-GLUCANASE	Glycosylase
25318	3162	TREHALOSE-6-PHOSPHATE PHOSPHATASE	Phosphatase
25326	3163	RECEPTOR-LIKE PROTEIN KINASE	Kinase, Protein
25352	3164	RECEPTOR-LIKE KINASE[PUTATIVE]	Kinase, Protein
25355	3165	LECTIN-LIKE PROTEIN KINASE[PUTATIVE]	Kinase, Protein
25357	3166	ANTHOCYANIDIN-3-GLUCOSIDE	Transferases
		RHAMNOSYLTRANSFERASE	
25370	3167	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
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25371	3168	RECEPTOR PROTEIN KINASE-LIKE PROTEIN	Kinase, Protein
25373	3169	XYLOGLUCAN ENDO-TRANSGLYCOSYLASE-LIKE	Glycosylase
		PROTEIN	<u> </u>
25383	3170	RECEPTOR PROTEIN KINASE-LIKE	Kinase, Protein
		PROTEIN[PUTATIVE]	
25386	3171	ARGININE METHYLTRANSFERASE[PUTATIVE]	Transferases
25389	3172	URIDYLYL TRANSFERASES-LIKE	Transferases
25399	3173	AMINO ACID TRANSPORTER PROTEIN-LIKE	Transporter
25406	3174	PROLYL 4-HYDROXYLASE, ALPHA SUBUNIT-LIKE	Hydroxylase
		PROTEIN	
25408	3175	PROTEIN PHOSPHATASE 2C-LIKE PROTEIN	Phosphatase
25412	3176	3-DEHYDROQUINATE SYNTHASE-LIKE PROTEIN	Synthase
25415	3177	ALPHA-MANNOSIDASE	Glycosylase
25419	3178	FERREDOXIN-NADP+ REDUCTASE	Reductase
25422	3179	CHALCONE ISOMERASE-LIKE PROTEIN	Isomerase
25433	3180	ELICITOR-INDUCIBLE RECEPTOR-LIKE	Receptor
		PROTEIN[PUTATIVE] .	:
25439	3181	PEROXIDASE (EMB CAA66964.1)	Oxidase
25443	3182	S-ADENOSYL-L-METHIONINE:SALICYLIC ACID	Transferases
		CARBOXYL METHYLTRANSFERASE-LIKE PROTEIN	
25446	3183	MANNAN ENDO-1,4-BETA-MANNOSIDASE	Glycosylase
25469	3184	UTP-GLUCOSE GLUCOSYLTRANSFERASE	Transferases
25479	3185	PROTEIN KINASE[PUTATIVE]	Kinase, Protein
25492	3186	PECTINESTERASE LIKE PROTEIN	Esterase
25503	3187	ZEAXANTHIN EPOXIDASE PRECURSOR	Oxidase
25509	3188	SUBTILISIN-TYPE PROTEASE-LIKE	Protease
25510	3189	DNA POLYMERASE ALPHA 1	Polymerase
25513	3190	MAP3K-LIKE PROTEIN KINASE[PUTATIVE]	Kinase, Protein
25515	3191	ANTHRANILATE N-	Transferases
		HYDROXYCINNAMOYL/BENZOYLTRANSFERASE-LIKE	
		PROTEIN	

25516	3192	ANTHRANILATE N-	Transferases
25510	3192	HYDROXYCINNAMOYL/BENZOYLTRANSFERASE-LIKE	Transiciases
		PROTEIN	
25500	0100		V
25520	3193	RECEPTOR PROTEIN KINASE-LIKE PROTEIN Kinase, Protein	
25523	3194	JDP-GLUCURONYLTRANSFERASE-LIKE PROTEIN Transfer	
25528	3195	RECEPTOR-LIKE PROTEIN KINASE Kinase, Protein	
25536	3196	CUCUMISIN-LIKE SERINE PROTEASE (GB AAC18851.1)	Protease
25540	3197	PEROXIDASE (EMB CAA66967.1) Oxidase	
25543	3198	N-ACETYLTRANSFERASE HOOKLESS1-LIKE PROTEIN	Transferases
113old	3199	ACCELERATED CELL DEATH 2; RED CHLOROPHYLL	Reductase
		CATABOLITE REDUCTASE [ARABIDOPSIS THALIANA]	
12old	3200	LETHAL LEAF-SPOT 1 HOMOLOG LLS1	Oxygenase
		[DIOXYGENASE DOMAIN][ARABIDOPSIS THALIANA].	
13288old	3201	(EC 4.2.99.8) CYSC1 [SIMILARITY] - ARABIDOPSIS	
		THALIANA.	
15402old	3202	CYSTEINE SYNTHASE (EC 4.2.99.8) 3A, CYTOSOLIC - Synthase	
		ARABIDOPSIS THALIANA.	
15792old	3203	CYSTEINE SYNTHASE (EC 4.2.99.8) ISOFORM 5-8,	Synthase
		CYTOSOLIC - ARABIDOPSIS THALIANA.	
15851old	3204	CYSTEINE SYNTHASE (EC 4.2.99.8) ACS1 -	Synthase
		ARABIDOPSIS THALIANA.	
1678old	3205	CYSTEINE SYNTHASE; O-ACETYLSERINE(THIOL)	Synthase
		LYASE [ARABIDOPSIS THALIANA].	
182old	3206	PROBABLE CYSTEINE SYNTHASE, CHLOROPLAST	Synthase
		PRECURSOR (O-ACETYLSERINE SULFHYDRYLASE) (O-	-
		ACETYLSERINE (THIOL)-LYASE) (CSASE) (OAS-TL)	
		(CS26).	
18927old	3207	PUTATIVE CYSTEINE SYNTHASE; 39489-37437	Synthase
		[ARABIDOPSIS THALIANA].	
203old	3208	5-METHYLTETRAHYDROPTEROYLTRIGLUTAMATE	Transferases
		HOMOCYSTEINE METHYLTRANSFERASE (VITAMIN-	
		B12-INDEPENDENT METHIONINE SYNTHASE	
		ISOZYME) (COBALAMIN-INDEPENDENT METHIONINE	
		SYNTHASE ISOZYME).	
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21308old	3209	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
21309old	3210	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
23001old	3211	4-DIPHOSPHOCYTIDYL-2C-METHYL-D-ERYTHRITOL	Synthase
		SYNTHASE [ARABIDOPSIS THALIANA].	
23094old	3212	CYSTEINE SYNTHASE, CHLOROPLAST PRECURSOR (O-	Synthase
		ACETYLSERINE SULFHYDRYLASE) (O-ACETYLSERINE	
		(THIOL)-LYASE) (CSASE).	
34209old	3213	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
34659old	3214	CYSTEINE SYNTHASE ATCYSC1 [ARABIDOPSIS Synthase	
		THALIANA].	
37280old	3215	CYSTEINE SYNTHASE, MITOCHONDRIAL PRECURSOR	Synthase
		(O-ACETYLSERINE SULFHYDRYLASE) (O-	
		ACETYLSERINE (THIOL)-LYASE) (CSASE).	
37284old	3216	CYSTEINE SYNTHASE (O-ACETYLSERINE	Synthase
		SULFHYDRYLASE) (O-ACETYLSERINE (THIOL)-LYASE)	
		(CSASE).	
39272old	3217	CHLOROPHYLL B SYNTHASE [ARABIDOPSIS	Synthase
		THALIANA].	
40108old	3218	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
40109old	3219	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA]. Synthase	
42762old	3220	5'-PHOSPHORIBOSYL-5-AMINOIMIDAZOLE	Synthase
		SYNTHETASE.	
42911old	3221	PUTATIVE CYSTEINE SYNTHASE [ARABIDOPSIS	Synthase
		THALIANA].	
44492old	3222	SIMILAR TO NICOTIANA 5-EPI-ARISTOLOCHENE	Synthase
		SYNTHASE (GB	
44907old	3223	CYSTEINE SYNTHASE (EC 4.2.99.8) 3A - ARABIDOPSIS	Synthase
		THALIANA.	
44988old	3224	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
45432old	3225	CYSTEINE SYNTHASE [ARABIDOPSIS THALIANA].	Synthase
46254old	3226	3-DEOXY-D-ARABINO-HEPTULOSONATE 7-	Synthase
		PHOSPHATE SYNTHASE.	
7417old	3227	2C-METHYL-D-ERYTHRITOL 2,4-CYCLODIPHOSPHATE	Synthase
		SYNTHASE [ARABIDOPSIS THALIANA].	

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- 233 -

Many annotations in publicly accessible data bases occur repeatedly, i.e. for various nucleic acid or amino acid sequences. The reasons for this are, to a minor extent, erroneous and/or redundant sequences and descriptions. To a major extent, this reflects the fact that proteins with the same function do indeed occur repeatedly in the genome. These different proteins can differ from each other for example by the regulation of their expression or by their cellular localization.

Many proteins belong to particular protein families. The skilled worker can draw conclusions with regard to the type of function, and thus also the possibility of an assay method for the polypeptide in question or its biological activity, from the protein family it belongs to. A description of such families of polypeptides and genes from Arabidopsis is obtainable for example in EP-A-1 033 405, but can also be found in the literature with which the skilled worker is familiar. Corresponding related information regarding the individual targets in Table 1 can be found in the document cited or in the general literature.

The analysis carried out for the purpose of the present invention, however, provides not only the general descriptions and the descriptions which are less suitable for the choice of herbicide targets in EP-A-1 033 405, but also the specificity of the polypeptide for the plant kingdom and the groups enzyme, receptor or channel (transporter) and more specific classes of these groups to which the proteins belong. The method according to the invention thus makes it possible to identify the particular suitability of a protein as target for finding lead structures for new herbicides exclusively with the aid of the method according to the invention. The classes which the polypeptides according to the invention were assigned to comprise, inter alia, acetylases, aldolases, amidases, amylases, anhydrases, arginases, ATPases, carboxylases, carrier-proteins, cellulases, channels, chelatases, chitinases, cyclases, deaminases, decarboxylases, dehydratases, dehydrogenases, desaturases, enolases, epimerases, esterases, furanases, furanosidases, galactosidases, galacturonases, glucanases, glucosidases, glucosylases, glucoronases, glycosylases, GTPases,

helicases, hydrolases, hydroxylases, isomerases, kinases, LACCases, lactonases, ligases, lipases, lyases, mannosidases, maturases, methylases, mutases, nucleases, nucleosidases, nucleotidases, oxidases, oxygenases, pectases, pectases, pectosidases, peptidases, permeases, phosphatases, phosphorylases, polymerases, proteases, racemases, receptors, reductases, sulfurylases, synthases, synthetases, transferases, transporters, transcriptases, xylanases and xylosidases.

The polypeptides which are identified by means of the method according to the invention are therefore particularly suitable as targets for finding new herbicidal active compounds. They are particularly suitable because they

a) have no homologous counterpart in animal organisms or in humans, according to the method according to the invention (determination of Evalues, alignment of data bases),

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b) were selected with a view that they are enzymes with small ligands or else receptors or channels which can, as a rule, be modulated, i.e. inhibited or activated, by small organic molecules or peptides and are therefore in principle open to being influenced by an active compound, and

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c) owing to the assignment to particular groups, make it possible for the skilled worker to select in a direct and obvious fashion assay methods which are suitable for the particular classes of polypeptides. To this end, the skilled worker can rely on the current literature or exploit the assay methods described in the present application.

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Subject-matter of the present invention is therefore furthermore the use of polypeptides found with the aid of the method according to the invention or of the nucleic acids encoding these polypeptides in methods for finding modulators of the polypeptides according to the invention or for finding new herbicidal compounds.

Subject-matter of the present invention is in particular the use of one of the polypeptides of SEQ ID NO: 1 to SEQ ID NO: 3227 in methods for finding modulators of these polypeptides or for finding new herbicidal compounds.

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The subject-matter of the present invention is furthermore the use of polypeptides which exert at least the biological activity of one of the polypeptides according to the invention and which encompass an amino acid sequence which has at least 60% identity, preferably 80% identity, especially preferably 90% identity, very especially preferably 97% identity, with a sequence of SEQ ID NO: 1 to SEQ ID NO: 3227 over its entire length in methods for finding modulators of the polypeptides or for finding new herbicidal active compounds.

The degree of identity of the amino acid sequences is determined for example with the aid of the program BLASTP + BEAUTY Version 2.0 4. (Altschul et al., 1997).

Preferred polypeptides which are used in the methods for finding modulators of the polypeptides according to the invention are those of SEQ ID NO: 1 to SEQ ID NO: 3227.

Based on the genetic code, a nucleic acid sequence encoding these polypeptides can be deduced in a simple fashion from the amino acid sequences of the polypeptides according to the invention, which amino acid sequences are shown in the sequence listing.

Such deduced nucleic acids can be used as probes and/or primers for detection and/or isolation of related polynucleotide sequences in different organisms, preferably in plants, through hybridization. Depending on the stringency of the conditions under which these probes and primers are used, polynucleotides exhibiting a wide range of similarity to those shown in Table 1 can be detected or isolated. "Stringency" as used herein is a function of probe length, probe composition (G/C content) and salt

concentration, organic solvent concentration and temperature of hybridization or wash conditions. Stringency is typically compared by the parameter  $T_m$ , which is the temperature of hybridization or wash conditions. Stringency is typically compared by the parameter  $T_m$  which is the temperature at which 50% of the complementary molecules in the hybridization are hybridized. High stringency conditions are e.g. those providing a condition of  $T_m$  5°C to 10°C. Medium or moderate stringency conditions are those providing  $T_m$  20°C to tm 29°C. Low stringency conditions are those providing for a condition of tm 40°C to  $T_m$  48°C. The relationship of hybridization conditions to  $T_m$  (in °C) is expressed in the following equation:

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$$T_m = 81.5 - 16.6 (\log_{10}[Na^+] + 0.41(\%G+C)) - (600/N),$$

where N is the length of the probe. This equation works well for probes comprising 14 to 70 nucleotides in length that are identical to the target sequence.

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Subject-matter of the present invention is therefore also the use of the nucleic acids encoding the polypeptides according to the invention in methods for finding new herbicidal compounds, and of DNA constructs which encompass one of the deduced nucleic acid sequences and a homologous or heterologous promoter.

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The term "homologous promoter" as used in the present context refers to a promoter which controls the expression of the gene in question in the original organism.

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The term "heterologous promoter" as used in the present context refers to a promoter which has properties other than the promoter which controls the expression of the gene in question in the original organism.

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The choice of heterologous promoters depends on whether pro- or eukaryotic cells or cell-free systems are used for expression. Examples of heterologous promoters are the cauliflower mosaic virus 35S promoter for plant cells, the alcohol dehydrogenase

promoter for yeast cells, the T3, T7 or SP6 promoters for prokaryotic cells or cell-free systems.

Subject-matter of the present invention is furthermore vectors comprising a nucleic acid encoding a polypeptide according to the invention or an abovementioned DNA construct. Vectors which can be used are all those phages, plasmids, phagemids, phasmides, cosmids, YACs, BACs, artificial chromosomes or particles which are suitable for particle bombardment, which are used in molecular biology laboratories.

10 Preferred vectors are pBIN (Bevan, 1984) and its derivatives for plant cells, pFL61 (Minet et al., 1992) or, for example, the p4XXprom. vector series(Mumberg et al.) for yeast cells, pSPORT vectors (Life Technologies) for bacterial cells, lambdaZAP (Stratagene) for phages or Gateway vectors (Life Technologies) for various expression systems in bacterial cells or Baculovirus.

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Subject-matter of the present invention is furthermore host cells comprising at least one nucleic acid encoding one of the polypeptides according to the invention or a DNA construct according to the invention or a vector according to the invention.

The term "host cell" as used in the present context refers to cells which do not naturally comprise the nucleic acids to be used in accordance with the invention.

Suitable host cells are prokaryotic cells, preferably *E. coli*, but also eukaryotic cells, such as cells of *Saccharomyces cerevisiae*, *Pichia pastoris*, insects, plants, frog oocytes and mammalian cell lines.

The term "polypeptides" as used in the present context refers not only to short amino acid chains which are usually termed peptides, oligopeptides or oligomers, but also to longer amino acid chains which are usually termed proteins. It encompasses amino acid chains which can be modified either by natural processes, such as post-

- 238 -

translational processing, or by chemical prior-art methods. Such modifications may occur at various sites and repeatedly in a polypeptide, such as, for example, on the peptide backbone, on the amino acid side chain, on the amino and/or the carboxyl terminal. For example, they encompass acetylations, acylations, ADP ribosylations, amidations, covalent linkages to flavins, haeme moieties, nucleotides or nucleotide derivatives, lipids or lipid derivatives or phosphatidylinositol, cyclisation, disulfide bridge formations, demethylations, cystine formations, formylations, gammacarboxylations, glycosylations, hydroxylations, iodinations, methylations, myristoylations, oxidations, proteolytic processings, phosphorylations, selenoylations and tRNA-mediated amino acid additions.

The polypeptides to be used in accordance with the invention may exist in the form of "mature" proteins or as parts of larger proteins, for example as fusion proteins. They can furthermore exhibit secretion or leader sequences, pro-sequences, sequences which make possible simple purification, such as polyhistidine residues, or additional stabilizing amino acids.

The polypeptides to be used in accordance with the invention need not constitute complete plant proteins but may also only be fragments thereof, as long as they retain at least one biological activity of the complete plant proteins. Polypeptides which exert the same type of biological activity as one of the proteins of Table 1 are still considered as being within the scope of the present invention. In this context, it is not necessary for the polypeptides to be used in accordance with the invention to be deducible from Arabidopsis proteins. Polypeptides which correspond to proteins of, for example, the plants given hereinbelow or fragments of these proteins which can still exert their biological activity are also considered as being within the scope of the present invention: tobacco, maize, wheat, barley, oats, oil seed rape, rice, rye, soya bean, tomatoes, legumes, potato plants, *Lactuca sativa*, Brassicae, woody species, *Physcomitrella patens*.

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In comparison with the corresponding regions of the naturally occurring polypeptides, the polypeptides according to the invention can have deletions or amino acid substitutions as long as they still exert at least one biological activity of the complete polypeptides. Conservative substitutions are preferred. Such conservative substitutions encompass variations, one amino acid being replaced by another amino acid from among the following group:

- 1. Small aliphatic residues, unpolar residues or residues of little polarity: Ala, Ser, Thr, Pro and Gly;
- 10 2. Polar, negatively charged residues and their amides: Asp, Asn, Glu and Gln;
  - 3. Polar, positively charged residues: His, Arg and Lys;
  - 4. Large aliphatic unpolar residues: Met, Leu, Ile, Val and Cys; and
  - 5. Aromatic residues: Phe, Tyr and Trp.
- 15 The following list shows preferred conservative substitutions:

Original residue	Substitution
Ala	Gly, Ser
Arg	Lys
Asn	Gln, His
Asp	Glu
Cys	Ser
Gln	Asn
Glu	Asp
Gly	Ala, Pro
His	Asn, Gln
Ile	Leu, Val, Met
Leu	Ile, Val, Met
Lys	Arg,
Met	Leu, Ile

- 240 -

Original residue	Substitution
Phe	Met, Leu, Tyr, Ile, Trp
Pro	Gly
Ser	Thr
Thr	Ser
Trp	Tyr, Phe
Tyr	Trp, Phe
Val	Ile, Leu

The skilled worker knows that the polypeptides of the present invention can be obtained by various routes, for example by chemical methods such as the solid-phase method. To obtain larger protein quantities, the use of recombinant methods is recommended. The expression of a cloned gene according to the invention or fragments thereof can be effected in a series of suitable host cells which are known to the skilled worker. To this end, a nucleic acid encoding one of the polypeptides according to the invention or a DNA construct according to the invention or vector is introduced into a host cell with the aid of known methods.

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The integration into the chromosome of the host cell, of the cloned nucleic acid according to the invention which is suitable for expressing the polypeptide according to the invention, is within the scope of the present invention. This nucleic acid or fragments thereof are preferably introduced into a plasmid, and the coding regions of the nucleic acids or fragments thereof are linked functionally to a constitutive or inducible promoter.

The basic steps for preparing the recombinant polypeptides according to the invention are:

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1. Obtaining a natural, synthetic or semi-synthetic nucleic acid (DNA) which encodes a polypeptide according to the invention.

- 241 -

PCT/EP01/09892

2. Introducing this DNA into an expression vector which is suitable for expressing the polypeptide according to the invention, either alone or as a fusion protein.

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- 3. Transforming a suitable host cell, preferably a prokaryotic host cell, with this expression vector.
- 4. Growing this transformed host cell in a manner which is suitable for expressing the polypeptide according to the invention. 10
  - 5. Harvesting the cells and isolating the polypeptide according to the invention by suitable, known methods.

15 In this context, the coding regions of the polypeptide according to the invention can be expressed for example in E. coli using the customary methods. Suitable expression systems for E. coli are commercially available, for example the expression vectors of the pET series, such as pET3a, pET23a, pET28a with His-tag or pET32a with His-tag for simple purification and thioredoxin fusion for increasing 20 the solubility of the expressed enzyme, and pGEX with glutathione synthetase fusion, and also the pSPORT vectors, with the possibility of transferring the coding region into different vectors of the Gateway system for various expression systems. The expression vectors are transformed into λ DE3-lysogenic E. coli strains, for example, BL21(DE3), HMS 174(DE3) or AD494(DE3). After the initial growth of the cells under standard conditions known to the skilled worker, expression is induced by means of IPTG. After induction of the cells, incubation is carried out for 3 to 24 hours at temperatures of from 18 to 37°C. The cells are disrupted by sonication in breaking buffer (10 to 200 mM sodium phosphate, 100 to 500 mM NaCl, pH 5 to 8. The protein expressed can be purified by chromatographic methods,

- 242 -

in the case of protein expressed with His-tag by chromatography on an Ni-NTA column.

Another favourable approach is the expression of a polypeptide according to the invention in commercially available yeast strains (for example, *Pichia pastoris*) or in insect cell cultures (for example Sf9 cells).

Alternatively, the polypeptides according to the invention can also be expressed in plants.

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A rapid method of isolating the polypeptides according to the invention which are synthesized by host cells using a nucleic acid encoding them starts with the expression of a fusion protein, it being possible for the fusion moiety to be affinity-purified in a simple manner. The fusion moiety can be, for example, glutathione S-transferase. The fusion protein can then be purified on a glutathione affinity column. The fusion moiety can be cleaved off by partial proteolytic cleavage for example at linkers between the fusion moiety and the polypeptide according to the invention which is to be purified. The linker can be designed such that it includes target amino acids, such as arginine and lysine residues, which define sites for trypsin cleavage. In order to generate such linkers, standard cloning methods using oligonucleotides may be applied.

Other purification methods which are possible are based on preparative electrophoresis, FPLC, HPLC (for example using gel filtration columns, reverse-phase columns or mildly hydrophobic columns), gel filtration, differential precipitation, ion-exchange chromatography and affinity chromatography.

The terms "isolation or purification" as used in the present context mean that the polypeptides according to the invention are separated from other proteins or other macromolecules of the cell or of the tissue. Preferably, a preparation comprising the

- 243 -

polypeptides according to the invention is at least 10-fold concentrated and especially preferably at least 100-fold concentrated with regard to the protein content over a host cell preparation.

The polypeptides according to the invention can also be affinity-purified without fusion moieties with the aid of antibodies which bind to the polypeptides.

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The polypeptides found here with the aid of the method according to the invention and the polypeptides which are homologous to them make possible the search for new specific herbicides; thus, ways are opened up of identifying lead structures, some of which may be completely new, with the aid of these targets. Thus, new interesting herbicides can be provided starting from such compounds which inhibit the present polypeptides.

Not only the enzymes, receptors and channels stated, but other proteins with other functions, too, can be filtered out for their plant specificity. This also applies to proteins whose function is as yet unknown.

Just as described above for finding new targets for herbicides, fungus- or insect-specific targets can be identified. For this purpose, the genomes of relevant phytopathogenic fungi, for example, *Magnaporthe* and many others, or insects, for example *Drosophila*, *Heliothis* and many others, are compared with the genomes of plants and animals. Thus, those enzymes, receptors and channels which are fungus-specific (and which do not occur in plants or animals) or which are insect-specific (and which do not occur in plants or higher animals, that is to say Chordata, in particular humans), can be identified.

The search for lead structures by target-based screening has played a key role for approximately 10 years in the search for pharmaceutical active compounds. In crop protection research, the same key position has emerged somewhat later. Owing to

- 244 -

this high relevance, a multiplicity of methods have been developed for verifying any new target. Also included are methods of expressing the genes in relevant systems with which the skilled worker in the field of various families of proteins or classes of enzymes is generally familiar.

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Enzymes and how they are affected by active compound candidate molecules can be measured quite generally on the basis of their enzymatic activity. The enzymatic conversion of starting materials to products can be determined in a multiplicity of ways: for example by monitoring the optical characteristics of the reaction solution (for example absorption, fluorescence, luminescence). If the enzymatic reaction cannot be monitored visually directly, the reaction can frequently be monitored by coupling with one or more further reactions, either enzymatic or non-enzymatic reactions, which can be monitored visually. As an alternative, a multiplicity of variants of binding assays have been developed which are based on measuring the binding of active compound candidate molecules to a protein. Binding assays can be carried out using radiolabeled or optically labeled detection molecules. Binding assays can also be carried out without labels, for example by methods of mass spectrometry or nuclear resonance spectrometry. This is in sharp contrast to the protein functions, which can be tested by cellular assays. Here, cells are constructed in a variety of ways which respond in a specific manner to the inhibition (or activation) of an enzyme (or receptor or channel). For example, bacteria can be constructed whose intrinsic enzyme has been switched off and was then replaced by a corresponding plant enzyme. When the action of active compound candidate molecules on the wild-type bacterial strain and the transgenic strain are compared, active compounds can be identified which relate to the plant enzyme. Cellular assays can preferably be used for assaying in particular receptors, but also channels. For example, non-plant cells can be constructed which recombinantly comprise a plant receptor and which visualize the response of the receptor to active compound candidate molecules visually. Thus, a luciferase can be expressed in receptormediated fashion, for example, and this luciferase can then be detected with high

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- 245 -

sensitivity. Channels which are ion-selective, in particular for calcium, can be detected for example by ion-selective stains.

The multiplicity of possibilities of opening up enzymes, receptors and channels to screening, preferably HTS or UHTS, is described in various reviews (see, for example, J. A. Landro et al., J. Pharmacol. Toxicol. Methods 44 (2201) 273 - 289). A large number of public fora exist for the specialists working in this field, such as, for example, the "Society for Biomolecular Screening" (Danbury, CT, USA) (www.sbsonline.org), which publishes its own periodical. The annual conferences of the "Society for Biomolecular Screening" reflect the current state of the art. It can therefore be said that it is currently possible to convert any desired protein into an HTS assay, it being possible for the difficulty or complexity of the assay method to vary, depending on the polypeptide.

Many assay systems whose aim it is to assay compounds and natural extracts are designed for high throughput numbers in order to maximize the number of substances studied within a given period. Assay systems which are based on cell-free procedures require purified or semipurified protein. They are suitable for a "first" assay, whose principal aim is to detect a potential effect of a substance on the target protein.

Effects such as cell toxicity are, as a rule, ignored in these *in vitro* systems. The assay systems test both inhibitory or suppressive effects of the substances and stimulatory effects. The efficacy of a substance can be tested by concentration-dependent test series. Control batches without test substances can be used for assessing the effects.

In the following text, methods shall be shown by way of example which can be exploited inter alia for finding modulators of the polypeptides according to the invention, the methods according to the invention including high-throughput screening (HTS) and ultra-high throughput screening (UHTS). Both host cells and

- 246 -

cell-free preparations comprising the nucleic acids according to the invention and/or the polypeptides according to the invention can be used for this purpose.

The examples given are understood as being a nonlimiting selection of methods which are possible for use for the purpose in accordance with the invention.

### Activity assays

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In order to find modulators of the polypeptides to be used according to the invention, for example a synthetic reaction mix (for example products of the in vitro transcription) or a cellular component, such as a crude cell extract, or any other preparation comprising the polypeptide to be used in accordance with the invention can be incubated together with one or more optionally labeled substrates or ligands of the polypeptides in the presence or absence of a candidate molecule, which may be an agonist or antagonist. The ability of the candidate molecule of increasing or inhibiting the activity of the polypeptide to be used in accordance with the invention can be seen from an increased or reduced conversion of the substrate. Molecules which lead to an increased activity of the polypeptides to be used in accordance with the invention are agonists. Molecules which lead to a reduction in the activity of the polypeptides to be used in accordance with the invention are probably inhibitors or antagonists. The detection of the biological activity of the polypeptides to be used in accordance with the invention can possibly be improved by what is known as a reporter system. Reporter systems as used herein comprise, but are not limited to, colorimetrically labeled substrates which are converted into a product, or a reporter gene which responds to changes in the activity or the expression of the polypeptides to be used in accordance with the invention.

## Binding assays

In order to find modulators of the polypeptides to be used according to the invention, for example a synthetic reaction mix (for example products of the *in vitro* transcription) or a cellular component, such as a crude cell extract, or any other

WO 02/10210

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- 247 -

PCT/EP01/09892

preparation comprising the polypeptide to be used in accordance with the invention can be incubated together with a labeled substrate or ligand of the polypeptides in the presence or absence of a candidate molecule, which may be an agonist or antagonist. The ability of the candidate molecule of increasing or inhibiting the activity of the polypeptide to be used in accordance with the invention can be seen from an increased or reduced binding of the labeled ligand. Molecules which bind well and lead to an increased activity of the polypeptides to be used in accordance with the invention are agonists. Molecules which bind well but do not trigger the biological activity of the polypeptides to be used in accordance with the invention are probably good antagonists. The detection of the biological activity of the polypeptides to be used in accordance with the invention can possibly be improved by what is known as a reporter system. Reporter systems as used herein comprise, but are not limited to, a reporter gene which responds to changes in the activity or expression of the polypeptides to be used in accordance with the invention, or other known binding assays.

#### Displacement assays

A further example of a method by means of which modulators of the polypeptides to be used in accordance with the invention can be found is a displacement assay in which the polypeptides to be used in accordance with the invention and a potential modulator are contacted under suitable conditions with a molecule which is known to bind to the polypeptides to be used in accordance with the invention, such as a natural substrate or ligand, or a substrate or ligand mimetic. The polypeptides to be used in accordance with the invention can be labeled themselves, for example radiolabeled or colorimetrically labeled, so that the number of the polypeptides which are bound to a ligand or which have undergone a conversion can be determined accurately. In this manner, the efficacy of an agonist or antagonist can be determined.

- 248 -

For the purposes of molecular interaction studies using a polypeptide according to the invention, or else with polypeptide variants which have been modified by in vitro mutagenesis or other known methods, a known analytical system may be employed, for example by Biacore AB, Uppsala, Sweden. In this system, (i) the polypeptide according to the invention or fragments thereof can be coupled to a biochip via known chemical methods (coupling via amines, thiols, aldehydes) or affinity binding (for example Streptavidin-Biotin, IMAC), or (ii) a ligand, for example a peptide or a small molecule, can be coupled to the chip. The binding, to the immobilized molecules, of a ligand in solution can be measured physically. In the case of the Biocore Instrument, the ligand is immobilized on a sensor chip with a thin gold layer. The solution of the analyte is perfused through a micro-flow cell on the chip. The binding of the analyte to the immobilized ligand increases the local concentration at the surface, the refractive index of the medium close to the gold layer gradually increasing. This affects the interaction between free electrons (plasmons) in the metal and photons which are emitted by the instrument. These physical changes are proportional to the mass and molecular number on the chip, the ligand-analyte binding is registered in real time, thus allowing the apparent association/dissociation rate to be determined (Fivash et al. 1998). Competition experiments validate the specificity of the binding. Analogous measurements also serve to determine the polypeptide domains are which are important for the binding of ligands, and to identify new, as yet unknown, ligands of the polypeptides according to the invention.

## Scintillation Proximity Assay (SPA)

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A possibility of identifying substances which modulate the activity of specific polypeptides according to the invention, such as, for example, receptor proteins, and polypeptides which are homologous thereto, is what is known as "Scintillation Proximity Assay" (SPA), see EP 015 473. This assay system exploits the interaction of a receptor with a radiolabeled ligand (for example a small organic molecule or a second radiolabeled protein molecule). The receptor is bound to microspheres or beads provided with scintillating molecules. As the radioactivity declines, the

WO 02/10210

- 249 -

PCT/EP01/09892

scintillating substance in the microsphere is excited by the subatomic particles of the radiolabel, and a detectable photon is emitted. The assay conditions are optimized in such a way that only those particles originating from the ligand lead to a signal which originate from a ligand bound to the receptor or to the polypeptide according to the invention.

In a possible embodiment, the polypeptide according to the invention is bound to the beads, either together with, or without, interacting or binding test substances. It would also be possible to use fragments of the polypeptides according to the invention. When a binding, for example radiolabeled, ligand binds to the immobilized polypeptide according to the invention, this ligand should inhibit or cancel out an existing interaction between the immobilized polypeptide according to the invention and the labeled ligand in order to bind itself in the contact area zone. Successful binding to the polypeptide according to the invention can then be detected by means of a flash of light. Analogously, an existing complex between an immobilized polypeptide and a free, labeled ligand is destroyed by the binding of a test substance, which leads to a drop in the intensity of the flash of light which is detected. In this case, the assay system corresponds to a complementary inhibition system.

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#### Two Hybrid System

An example of an assay system based on intact cells is what is known as the Two Hybrid System, which is particularly suitable for those polypeptides which have a suitable interaction partner in the cell - a further polypeptide or peptide. A specific example is what is known as the interaction trap. This is a genetic selection of interacting proteins in yeast (see, for example, Gyuris et *al.* 1993). The assay system is designed to detect and describe the interaction of two proteins, owing to an interaction which has taken place leading to a detectable signal.

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- 250 -

Such an assay system can also be adapted to the testing of large numbers of test substances in a given period.

The system is based on the construction of two vectors, the bait vector and the prey vector. A gene encoding a polypeptide according to the invention or fragments thereof is cloned into the bait vector and then expressed as fusion protein together with the LexA protein, a DNA binding protein. A second gene encoding an interaction partner of the polypeptide in question is cloned into the prey vector, where it is expressed as fusion protein together with the B42 prey protein. Both vectors are present in a *Saccharomyces cerevisiae* host which contains copies of LexA-binding DNA 5' of a lacZ or HIS3 reporter gene. If an interaction takes place between the two fusion proteins, activation of the transcription of the reporter gene results. If the presence of a test substance results in inhibition or interference with the interaction, the two fusion proteins can no longer interact and the product of the reporter gene is no longer produced.

- 251 -

# Calcium Imaging

Calcium imaging or signalling must be considered as a further method of detecting substances which interact with polypeptides according to the invention. This method is suitable, for example, for receptors which act as  $Ca^{2+}$  channels. Here, calcium indicators are employed with the aid of which changes in the intracellular calcium level are made detectable. Within the scope of these methods, cells which express the relevant polypeptide according to the invention are employed, and these cells are loaded with calcium indicators. Upon UV excitation, an influx of calcium caused by an HC110-R agonist, or the release of intracellular calcium, leads to a change in absorption as a function of the calcium load of the indicator. In such a system, an antagonist can be recognized by the complete or partial suppression of the calcium signal induced by the agonist (for example  $\alpha$ -LTX). Suitable calcium indicators which are possible for this purpose are Fura-2 (Sigma) or Indo-1 (Molecular Probes).

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Further calcium indicators can be excited by visible light and change their fluorescence behaviour detectably as a function of their calcium load. The indicators Fluo-3 and Fluo-4 show high affinity for calcium. Fluo-4, which has the stronger fluorescence signal, is particularly suitable for measurements in test systems where the cells are employed only at low density, as is the case for HEK293 cells. Further indicators are Rhod-2, x-Rhod-1, Fluo-5N, Fluo-5F, Mag-Fluo-4, Rhod-5F, Rhod-5N, Y-Rhod-5N, Mag-Rhod-2, Mag-X-Rhod-1, Calcium Green-1 and -2, Calcium Green-5N, Oregon Green 488 BAPTA-1, Oregon Green 488 BAPTA-2 and -5N, Fura Red, Calcein and the like.

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An alternative to loading cells with calcium indicators is the recombinant expression of photoproteins in the target cells. Once these photoproteins have formed a complex with calcium ions, they react in the form of a light emission. A photoprotein which has already been used often in a large number of studies and assay systems is aequorin. In this assay method, the cells which simultaneously express the target

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protein and the aequorin are first loaded with the luminophore coelenterazin. The apoaequorin formed by the cells forms a complex with the coelenterazin and carbon dioxide. If calcium subsequently enters the cell and binds to the complex, carbon dioxide and blue light are emitted (emission maximum ~466 nm). The light emission correlates with the calcium concentration which prevails intracellularly.

Subject-matter of the present invention is therefore in particular also the use of the polypeptides of the Table 1 which have been identified with the aid of the present method in methods of finding modulators of the polypeptides according to the invention.

Subject-matter of the present invention is furthermore the use of nucleic acids encoding these plant proteins, DNA constructs comprising them, host cells comprising them, or antibodies which bind to these proteins in methods of finding modulators of the polypeptides according to the invention.

The term "agonist" as used in the present context refers to a molecule which accelerates or increases the activity of the protein.

The term "antagonist" as used in the present context refers to a molecule which slows down or prevents the activity of the protein.

The term "modulator" as used in the present context constitutes the generic term for agonist and antagonist. Modulators can be small organochemical molecules, peptides or antibodies which bind to the polypeptides to be used in accordance with the invention. Furthermore, modulators can be small organochemical molecules, peptides or antibodies which bind to a molecule which, in turn, binds to the polypeptides to be used in accordance with the invention, thus influencing their biological activity. Modulators can constitute natural substrates and ligands or of

- 253 -

PCT/EP01/09892

structural or functional mimetics thereof. However, the term "modulator" does not extend to the natural substrates and to ATP.

The modulators are preferably small organochemical compounds.

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WO 02/10210

The binding of the modulators to the proteins to be used in accordance with the invention can modify the cellular processes in such a way which lead to the death of the plants treated therewith.

Subject-matter of the present invention are therefore also modulators which have been found with the aid of one of the polypeptides described in accordance with SEQ ID NO:1 to SEQ ID NO:3227 for identifying modulators of a polypeptide.

Subject-matter of the invention is furthermore the use of modulators of the polypeptides in accordance with SEQ ID NO:1 to SEQ ID NO:3227 as herbicides.

Furthermore, the present invention comprises methods of finding chemical compounds which modify the expression of the polypeptides to be used in accordance with the invention. Such "expression modulators", again, can constitute growth-regulatory or herbicidal active compounds. Expression modulators can be small organochemical molecules, peptides or antibodies which bind to the regulatory regions of the nucleic acids encoding the polypeptides which are to be used in accordance with the invention. Furthermore, expression modulators can be small organochemical molecules, peptides or antibodies which bind to a molecule which, in turn, binds to regulatory regions of the nucleic acids encoding the polypeptides to be used in accordance with the invention, thus influencing their expression. Expression modulators can also be antisense molecules.

- 254 -

The present invention therefore also extends to the use of modulators of the polypeptides according to the invention or of expression modulators of same as plant growth regulators or herbicides.

Subject-matter of the present invention are also expression modulators of proteins which are found with the aid of any above-described method of identifying expression modulators of the proteins.

Subject-matter of the invention is also the use of expression modulators as herbicides.

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- 257 -

PCT/EP01/09892

# **Patent Claims**

**WO** 02/10210

1. Method of identifying target proteins for herbicidally active compounds, comprising the following steps:

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alignment of a nucleic acid sequence or amino acid sequence (Group 1 a) sequence) from plants with a nucleic acid sequence or amino acid sequence from non-plant organisms (group 2 sequence) using suitable search parameters,

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b) determination of the E-value of the group 1 sequence and a similar group 2 sequence, and

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selecting group 1 sequences in which the exponent of the E-value c) exceeds that of the most similar group 2 sequence at least by a factor of 3.

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2. Method according to Claim 1, characterized in that, in a further step, those group 1 sequences are selected which are essential for the plant and, if appropriate, naturally have small ligands.

3. Method according to Claim 1 or 2, characterized in that the E-value is not lower than 10<sup>-30</sup>.

25 Use of polypeptides or of nucleic acids encoding them which are found in a 4. method according to one of Claims 1 to 3 in a method of identifying modulators of these polypeptides or nucleic acids.

- 258 -

- Use of one of the polypeptides in accordance with SEQ ID NO: 1 to SEQ ID NO: 3227 and of the nucleic acids encoding them in methods of identifying modulators of these polypeptides.
- Method of finding a chemical compound which modulates the activity of one of the polypeptides in accordance with SEQ ID NO: 1 to SEQ ID NO: 3227, comprising the following steps:
- (a) contacting a preparation or host cell comprising the polypeptide with a chemical compound or a mixture of chemical compounds under conditions which permit the interaction of a chemical compound with the polypeptide, and

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- (b) identifying the chemical compound which specifically influences the activity of the polypeptide.
- 7. Method of finding a chemical compound which binds to one of the polypeptides in accordance with SEQ ID NO: 1 to SEQ ID NO: 3227 and/or which displaces a natural substrate or a natural ligand, comprising the following steps:
  - (a) contacting a preparation or host cell comprising the polypeptide with a chemical compound or a mixture of chemical compounds under conditions which permit the interaction of a chemical compound with the polypeptide, and
  - (b) identifying the chemical compound which specifically binds to the polypeptide, and/or

- 259 -

(c) identifying the chemical compound which specifically displaces a natural substrate or a natural ligand.

8. Method of finding a chemical compound which modulates the cellular function of one of the polypeptides in accordance with SEQ ID NO: 1 to SEQ ID NO: 3227, comprising the following steps:

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- (a) contacting a host cell which expresses the polypeptide with a chemical compound or a mixture of chemical compounds under conditions which permit the interaction of the chemical compound with the cell and/or the polypeptide, and
- (b) identifying the chemical compound which specifically influences the cellular function of the polypeptide.
- 9. Method of finding a compound which modifies the expression of the polypeptide in accordance with SEQ ID NO: 1 to SEQ ID NO: 3227, comprising the following steps:
- 20 (a) contacting a host cell expressing the polypeptide with a chemical compound or a mixture of chemical compounds,
  - (b) determining the polypeptide concentration, and
- 25 (c) identifying the compound which specifically influences the expression of the polypeptide.
  - Use of a modulator of one of the polypeptides in accordance with SEQ IDNO: 1 to SEQ ID NO: 3227 as herbicide.

- 260 -

11. Herbicides which are found in a method according to Claim 6 or 7.